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Dust temperature and time-dependent effects in the chemistry of photodissociation regions

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ABSTRACT

When studying chemistry of photodissociation regions (PDRs), time dependence becomes important as visual extinction increases, since certain chemical time-scales are comparable to the cloud lifetime. Dust temperature is also a key factor, since it significantly influences gas temperature and mobility on dust grains, determining the chemistry occurring on grain surfaces. We present a study of the dust temperature impact and time effects on the chemistry of different PDRs, using an updated version of the Meijerink PDR code and combining it with the time-dependent code Nahoon. We find the largest temperature effects in the inner regions of high G_0 PDRs, where high dust temperatures favour the formation of simple oxygen-bearing molecules (especially that of O_2), while the formation of complex organic molecules is much more efficient at low dust temperatures. We also find that time-dependent effects strongly depend on the PDR type, since long time-scales promote the destruction of oxygen-bearing molecules in the inner parts of low G_0 PDRs, while favouring their formation and that of carbon-bearing molecules in high G_0 PDRs. From the chemical evolution, we also conclude that, in dense PDRs, CO_2 is a late-forming ice compared to water ice, and confirm a layered ice structure on dust grains, with H_2O in lower layers than CO_2 . Regarding steady state, the PDR edge reaches chemical equilibrium at early times ($\lesssim 10^5$ yr). This time is even shorter ($< 10^4$ yr) for high G_0 PDRs. By contrast, inner regions reach equilibrium much later, especially low G_0 PDRs, where steady state is reached at $\sim 10^6$ – 10^7 yr.

Key words: astrochemistry – ISM: abundances – ISM: clouds – photodissociation region (PDR).

1 INTRODUCTION

Photodissociation regions (PDRs) are characterized by their exposure to strong far-ultraviolet (FUV) radiation fields ($6 < h\nu < 13.6$ eV), which results in the heating of gas up to relatively high temperatures. These regions are important from a chemical point of view, since they play a key role in the formation of new species along the molecular cloud, as the UV radiation penetrates into the region.

PDRs can be found in different environments of the Milky Way, such as in massive star-forming regions (e.g. Tauber et al. 1994; Hora et al. 2004), close to cooler stars which emit enough FUV radiation to form lower density and lower excitation PDRs (e.g. Wyrkowski et al. 2000; Köhler et al. 2014), on the surface of

protoplanetary discs (e.g. van Dishoeck, Jonkheid & van Hemert 2006; Bergin et al. 2007), at the edge of molecular clouds (e.g. Spezzano et al. 2016), and also near evolved stars which emit strong FUV radiation (Meixner et al. 2001). This is also the case of the PDRs detected in planetary nebulae (PNe) through the emission of atomic fine structure lines, e.g. the ground state line of [C I] in NGC 6720 and in NGC 7293 (Bachiller et al. 1994; Young 1997). PDRs are also responsible for most of the non-stellar infrared emission from galaxies (e.g. Fuente et al. 2008; Bayet et al. 2009).

The large variety of environments where PDRs are found determines their physical conditions. In particular, PDRs can be diffuse, with gas density $n \sim 10$ – 10^2 cm $^{-3}$, or dense, with $n > 10^4$ cm $^{-3}$, while the incident FUV flux may range from the interstellar radiation field (ISRF) to 10^6 times the ISRF in the surroundings of an O star. PDRs are characterized by a layered structure, as a result of the interaction of the radiation with the gas and dust. Typically, they

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contain an outer layer (the edge of the cloud where visual extinction is $A_V < 1$ mag) of partially ionized gas, where hydrogen is atomic and carbon is predominantly in the form of C^+ . The transition to molecular hydrogen occurs in a region where carbon is still ionized, while the neutral carbon layer and the transition to CO occur where hydrogen is already fully molecular (e.g. Tielens & Hollenbach et al. 1985; Joblin et al. 2018).

Ultraviolet photons dominating the energy balance of PDRs do not only influence significantly their chemical structure, but also the time evolution of the interstellar medium (ISM) conditions regulating the star formation processes. At low visual extinctions, physical and chemical processes dominated by interactions with photons are fast compared to dynamical processes. However, at large visual extinctions, certain chemical time-scales are comparable to cloud lifetimes and time dependence becomes a key factor in the study of PDRs.

There are several authors (e.g. Bertoldi & Draine 1996; Kirsanova, Wiebe & Sobolev 2009; Morata & Herbst 2008; Hollenbach et al. 2009; Motoyama et al. 2015; Le Gal et al. 2017) who include time dependence in their PDR codes, however they do consider a simpler treatment of surface chemistry than in this study. In Esplugues et al. (2016), we showed not only the effects of varying the density and the intensity of the radiation field on the chemical evolution of different PDRs, but also the importance of considering surface chemistry when studying the chemical structure of molecular clouds exposed to different UV radiation fields. We derived that some parameters (such as the type of grain substrate and the probability of desorption) can alter the chemistry occurring on grain surfaces, leading to significant differences in the abundances of gas-phase species. Esplugues et al. (2016) also showed that many of these differences become even larger as the visual extinction increases, making evident the need of considering time dependence.

In this paper, we focus on time dependence and its effects on the chemical evolution of different PDRs, as well as on the role of dust temperature (T_{dust}) in the PDR chemistry. We carry out this study using an updated version of the Meijerink PDR code (presented in Section 2) with new solid species and surface chemical reactions, as well as with a new way to calculate the chemical desorption¹ probabilities for two-body reactions. In Section 3, we present the temperature study considering two different expressions for T_{dust} . In Section 4, we combine our steady-state code with the time-dependent code Nahoon to analyse the chemical evolution as a function of time and visual extinction. Section 5 contains the discussion of results, and a comparison with observations. In addition, we provide results for the time at which steady state is reached in each PDR type. A summary of the main conclusions is presented in Section 6.

2 THE STEADY-STATE PDR CODE

2.1 Gas chemistry

The updated Meijerink PDR code consists of 7503 chemical gas-phase reactions from the Kinetic Database for Astrochemistry

¹Chemical desorption process occurs when there is excess energy after the two-body reaction on dust grains. In order to desorb, the newly formed molecule has to convert a fraction of this excess formation energy into kinetic energy and, in particular, into motion perpendicular to the substrate (Minissale, Congiu & Dulieu 2014; Minissale et al. 2016).

Table 1. Solid species in our PDR code.

H	H ₂	HCO	C
H _c	HO ₂	H ₂ CO	CH
O	H ₂ O	CH ₃ O	CH ₂
O ₂	H ₂ O ₂	CH ₃ OH	CH ₃
O ₃	CO	N	CH ₄
OH	CO ₂	N ₂	S

H_c refers to the strong interaction between hydrogen and the grain surface (chemisorption), where the forces involved are similar to valence forces (see Cazaux & Tielens 2002 for more details).

(KIDA, Wakelam et al. 2012).² They include bimolecular reactions, charge-exchange reactions, radiative associations, associative detachment, dissociative recombination, neutralization reactions, ion-neutral reactions, ionization or dissociation of neutral species by UV photons, and ionization or dissociation of species by direct collision with cosmic ray particles or by secondary UV photons following H₂ excitation.

The heating mechanisms considered in the thermal balance of the code are photoelectric effect on grains, carbon ionization heating, H₂ photodissociation heating by UV photons, H₂ collisional de-excitation heating, gas-grain collisional heating, gas-grain viscous heating, and cosmic ray heating. As cooling mechanisms, we consider fine-structure line cooling (being [C II] at 158 μm and [O I] at 63 μm and at 146 μm the most prominent cooling lines), metastable-line cooling (including lines of C, C⁺, Si, Si⁺, O, O⁺, S, S⁺, Fe, and Fe⁺), recombination cooling, and molecular cooling by H₂, CO, and H₂O (see Meijerink & Spaans 2005 and Esplugues et al. 2016 for more details).

2.2 Dust chemistry

In a precedent study, we updated the Meijerink PDR core with the chemistry occurring on grain surfaces and added 18 solid species. In this study, we have included six additional solid species: S, C, CH, CH₂, CH₃, and CH₄ (see all the solid species considered in Table 1). We have also updated the surface chemical network implemented in the Meijerink code taking recent laboratory experiments (e.g. Dulieu et al. 2013; Minissale et al. 2015, 2016) into account. The surface processes considered in the code are adsorption, thermal desorption, chemical desorption, two-body reactions, photo processes, and cosmic ray processes. All these processes are described in detail in Esplugues et al. (2016). The other main change introduced in this new version of the code is the way to calculate the chemical desorption probabilities for two-body reactions in order to take more scenarios for the formation of chemical products into account. In particular, in the previous version of the Meijerink code, given the surface chemical reaction $JA + JB \rightarrow JC + JD$ (where Ji means solid i), we considered only two possibilities based on an empirical physical model adjusted on experimental data:



and



In this new version of the Meijerink code, however, we propose one way to extend it by considering chemical desorption per product,

²<http://kida.obs.u-bordeaux1.fr>.

which implies four possibilities:



and



where the chemical desorption coefficients CD_{JC} and CD_{JD} of the species JC and JD, respectively, are independent and calculated using:

$$\text{CD} = \exp\left(\frac{-E_{\text{binding}}}{\epsilon \Delta H_{\text{R}}/N}\right). \quad (7)$$

The factor E_{binding} is the binding energy of the desorbed product [$E_{\text{binding}}(\text{JC})$ for the case of CD_{JC} and $E_{\text{binding}}(\text{JD})$ for the case of CD_{JD} , using values shown in Table A3], $\epsilon \Delta H_{\text{R}}/N$ represents the total chemical energy available for the kinetic energy perpendicular to the grain surface, ΔH_{R} being the reaction enthalpy, $N = 3 \times n_{\text{atoms}}$ is the degree of freedom considering the atoms of the two newly formed molecules, and ϵ the fraction of kinetic energy retained by the product of mass m colliding with the surface, which has an effective mass M (see Cazaux et al. 2016; Minissale et al. 2016 for more details):

$$\epsilon = \frac{(M - m)^2}{(M + m)^2}. \quad (8)$$

The desorption probabilities for the four chemical reactions are:

$$\delta_{\text{R1}} = 100 - \max(\text{CD}_{\text{JC}}, \text{CD}_{\text{JD}}), \quad (9)$$

$$\delta_{\text{R2}} = \min(\text{CD}_{\text{JC}}, \text{CD}_{\text{JD}}), \quad (10)$$

$$\delta_{\text{R3}} = \text{CD}_{\text{JD}} - \min(\text{CD}_{\text{JC}}, \text{CD}_{\text{JD}}), \quad (11)$$

and

$$\delta_{\text{R4}} = \text{CD}_{\text{JC}} - \min(\text{CD}_{\text{JC}}, \text{CD}_{\text{JD}}), \quad (12)$$

where $\delta_{\text{R1}} + \delta_{\text{R2}} + \delta_{\text{R3}} + \delta_{\text{R4}} = 100$ per cent. In this case, unlike Esplugues et al. (2016), we calculate the desorption probabilities for each reaction using the binding energies of both products. This new approach considers therefore the fact that C and D are different products, with different energies and different degrees of freedom, and that, in exothermic reactions, the energy released is dissipated in a different manner for C and D. In addition, this formulation also reproduces the experimental results where only one product is observed, even if the considered reactions would have two products. See Appendix A for the list of chemical reactions occurring on grain surfaces that are included in the Meijerink PDR code.

3 DUST TEMPERATURE

Interstellar dust is a ubiquitous component of the ISM, whose mass is only about 0.7 per cent of the gas (Fisher et al. 2014). In spite of this low value, dust grains have an important impact on the chemistry and thermodynamics of molecular clouds. In particular,

the temperature of dust grains influences the gas temperature through heating and cooling processes along with chemical reaction rates. In addition, dust grain surfaces are also powerful interstellar catalysts since they are responsible for most of the production of the simplest (H_2) to the most complex (pre-biotic) molecules observed in the Universe.

Several analytical expressions for the dust temperature can be found in the literature, such as those from Hollenbach, Takahashi & Tielens (1991), Zucconi, Walmsley & Galli (2001), and Garrod & Pauly (2011). These expressions are calculated in different ways. The solution by Hollenbach et al. (1991) assumes a one-sided slab geometry and combines the heating by ultraviolet (UV) photons, cosmic microwave background (CMB), and the re-processed infrared (IR). The derived temperature is a function of the intensity of the radiation field (G_0) and of the visual extinction (A_V), although the A_V dependence only takes into account the attenuation of UV photons. The expression provided by Zucconi et al. (2001) considers the contributions from the visual/near-infrared, mid-infrared, and far-infrared (FIR), and the dust temperature solution is given for the range $10 \lesssim A_V \lesssim 400$ mag. This expression is based on the observed dust temperature of L1544 at various A_V and it is only a function of the visual extinction. To obtain it, the authors solve the thermal balance without considering the UV field. They only include the visual and infrared part of the spectrum. The dust temperature expression provided by Garrod & Pauly (2011) was designed for low A_V regions and to be combined with that from Zucconi et al. (2001) for larger extinctions. This expression is only a function of A_V .

A recent analytical expression for the dust temperature (T_{dust}) has been determined by Hocuk et al. (2017) from first principles for dust in thermal equilibrium by considering in detail the ISRF, the attenuation of radiation, the dust opacities, and various grain material compositions (graphite, silicates SiO_2 and MgFeSiO , and carbonaceous silicate mixtures). This expression is:

$$T_{\text{dust}} = [11 + 5.7 \times \tanh(0.61 - \log_{10}(A_V))] \times (\chi^{1/5.9}), \quad (13)$$

with χ the intensity of the radiation field in Draine units³. The final solutions were compared with those obtained from the Monte Carlo radiative transfer code RADMC-3D⁴ and with observational results from several interstellar regions observed with Herschel. See Hocuk et al. (2017) for more details.

Depending on the size of dust grains, their temperature can present significant variations on short time-scales (seconds to minutes) as derived by Cuppen, Morata & Herbst (2006) and Iqbal, Acharyya & Herbst (2014) using Monte Carlo simulations, and by Bron, Le Boulout & Le Petit (2014) using an analytical approach. In particular, the smallest grains (radii $a \lesssim 50$ Å) undergo very large temperature fluctuations (more than 30 K). These variations are equivalent to consider PDRs with radiation intensities of two different orders of magnitude, which significantly varies the chemistry (see Fig. 1 and sections below). Therefore, in the case of very small grains, it is not realistic to consider an average temperature. However, larger dust grains (and especially those with a size $a \geq 200$ Å) can be approximated as having a steady temperature (Draine & Li 2001), since their temperature fluctuations are lower than 3 K (Cuppen et al. 2006). Nevertheless, it should be noted that recent studies (Pauly & Garrod 2016) show that the dust temperature choice is far from being trivial, since other factors, such as the mantle

³Draine field $\simeq 1.7 \times$ Habing field (Habing 1968, Draine 1978).

⁴<http://www.ita.uni-heidelberg.de/~dullemond/software/radmc-3d>.

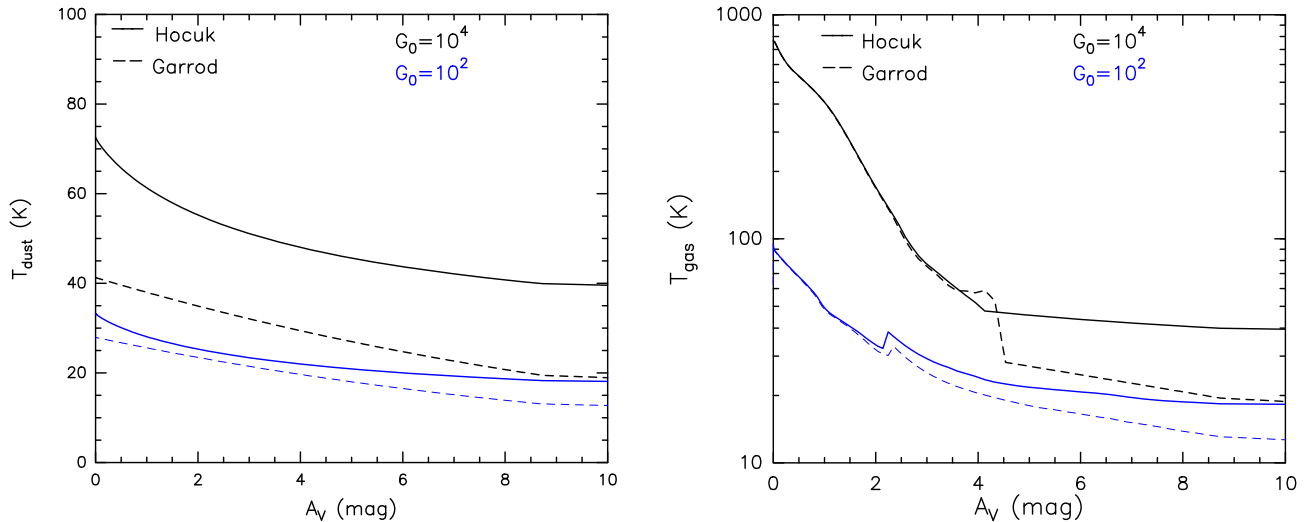


Figure 1. Dust (left) and gas (right) temperature for two PDR models with $G_0 = 10^2$ and $n = 10^5 \text{ cm}^{-3}$ (blue lines), and with $G_0 = 10^4$ and $n = 10^5 \text{ cm}^{-3}$ (black lines), considering analytical expressions from Garrod & Pauly (2011) (dashed line) and from Hocuk et al. (2017) (solid line).

growth and its time evolution, can also vary the dust temperature. In particular, they find dust temperature variations of $\sim 11 \text{ K}$ for grains with $a \lesssim 0.01 \mu\text{m}$, while the temperature variation is only $\lesssim 5 \text{ K}$ for grains with $a \gtrsim 0.1 \mu\text{m}$. In any case, these results make also evident the fact that the larger the grain sizes, the lower the dust temperature variations. Considering this fact and in order to avoid large local dust temperature fluctuations in short time-scales that could significantly alter the chemistry when studying the effects of other parameters (e.g. the effect of increasing the radiation field intensity), we have assumed an MRN grain size distribution (Mathis, Rumpl & Nordsieck 1977) in the Meijerink PDR code, with grain radius limited to $50 \text{ \AA} < a < 0.25 \mu\text{m}$, for which it is reasonable to consider an average dust grain temperature.

In Esplugues et al. (2016), we calculated T_{dust} through the expression from Garrod & Pauly (2011).⁵ Here, we also consider in our analysis the T_{dust} expression from Hocuk et al. (2017). Fig. 1 shows the dust temperature values from these two expressions for two PDRs with different intensity of radiation field⁶ ($G_0 = 10^2$ and $G_0 = 10^4$) in the interval $0 \leq A_V \leq 10 \text{ mag}$. For the PDR with the lowest G_0 (blue), the differences for T_{dust} between both expressions are lower than 10 K . However, for the most extreme PDR (black), these differences are of up to 30 K , leading to significant differences in the chemistry of the considered regions at intermediate and large visual extinctions (see Fig. 2). In Section 5.1, we analyse in detail the impact of considering both dust temperature expressions on the chemistry of several molecule families.

4 TIME DEPENDENCE

In a molecular cloud, as the visual extinction increases ($A_V > 1 \text{ mag}$), certain chemical time-scales become comparable to cloud lifetimes (10^6 – 10^7 yr) and steady-state chemistry does not apply. In

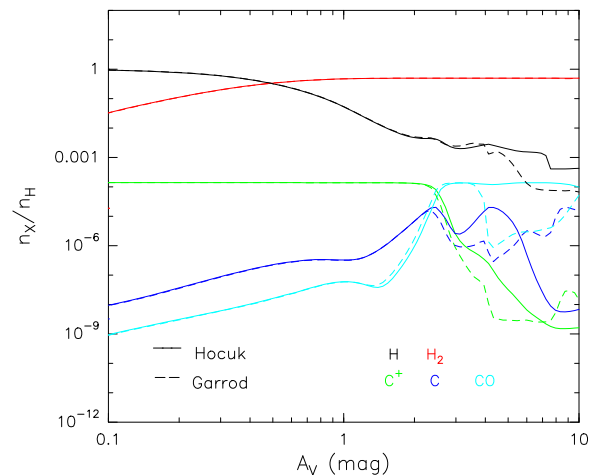


Figure 2. Abundances of H, H_2 , C^+ , C, and CO obtained with the updated Meijerink PDR code using T_{dust} from Hocuk et al. (2017) (solid lines) and from Garrod & Pauly (2011) (dashed lines). Results are obtained considering $G_0 = 10^4$ and $n = 10^5 \text{ cm}^{-3}$.

these cases, time-dependent solutions to the chemistry are therefore needed. This is the case of PDRs. At low visual extinctions ($A_V \lesssim 1 \text{ mag}$), the energy balance is dominated by FUV photons and the chemical time-scales are very short ($\lesssim 10^5 \text{ yr}$) compared to the molecular cloud lifetime. However, in the opaque interiors of the cloud ($A_V > 6 \text{ mag}$), the chemistry is dominated by a low FUV flux and by long chemical time-scales (e.g. the corresponding time-scale to cosmic ray desorption of CO ice is from 3×10^5 to $3 \times 10^9 \text{ yr}$, depending on the assumptions regarding the CO desorption process, Hollenbach et al. 2009). At intermediate depths, UVs are attenuated by dust extinction, but photodesorption still prevents total freeze-out.

To study the effects of time dependence on the chemistry of PDRs, we have coupled the Meijerink PDR code with the time-dependent code Nahoon. In this way, the PDR code provides a fixed physical structure (density, temperature) and we perform post-processing computing by calculating the time-dependent chemistry of the medium with Nahoon. Grains are initially bare and the

⁵Dust temperature expression derived from Garrod & Pauly (2011), but with an adaptation to include dependence with the intensity of the radiation field [Garrod (private communication), see Esplugues et al. 2016 for more details].

⁶We use G_0 , the Habing field (Habing 1968), as the normalization in which we express the incident FUV radiation field, where $G_0 = 1$ corresponds to a flux of $1.6 \times 10^{-3} \text{ erg cm}^{-2} \text{ s}^{-1}$.

Table 2. Adopted model parameters in our PDR code.

Model	G_0	n_{H} (cm^{-3})
1	10^2	10^5
2	10^4	10^5
3	10^4	10^6

formation of ices takes place during the evolution of the interstellar gas cloud, starting from a diffuse, fully atomic stage to a molecular phase illuminated and warmed up by a nearby star. We follow the composition at any time with chemical network using rate equations that incorporates grain surface reactions on two different substrates (bare and icy grains). The chemistry evolves over a period of 10^7 yr. The Nahoon code has been modified to have the same chemical network and chemical processes as those included in the Meijerink PDR code. In particular, to the gas-phase chemistry network provided by KIDA, we added our grain surface chemistry network as detailed in Section 2.2. The grain surface processes taken into account are identical to those used in the PDR code: adsorption, thermal desorption, two-body reactions, chemical desorption, desorption by UV photons and cosmic rays, and dissociation by UV photons and cosmic-ray-induced UV photons. A more detailed description of Nahoon, which is publicly available on KIDA, can be found in Wakelam et al. (2012). In Section 5.2, we analyse the time effects on the chemical evolution of different PDR types.

5 RESULTS AND DISCUSSION

We show the results for several molecule families through three different PDR models: with density $n = 10^5 \text{ cm}^{-3}$ and $G_0 = 10^2$ (Model 1), with $n = 10^5 \text{ cm}^{-3}$ and $G_0 = 10^4$ (Model 2), and with $n = 10^6 \text{ cm}^{-3}$ and $G_0 = 10^4$ (Model 3), see Table 2. These models have been chosen to analyse how the dust temperature and time-dependent effects vary depending on the type of PDR.

5.1 Dust temperature effects

Figs 3–5 show abundances for several species obtained with the most recent version of the Meijerink PDR code presented here, considering T_{dust} from Hocuk et al. (2017) (green dashed lines) and from Garrod & Pauly (2011) (blue dotted lines). We obtain that the chemical impact of considering different dust temperature significantly varies depending on the characteristics of the PDR, the visual extinction range, and the type of molecule. Below we analyse the dust temperature effects considering several molecule families.

5.1.1 Simple oxygen-bearing molecules

Fig. 3 shows the abundances of simple oxygen-bearing species (OH, O_2 , and H_2O) considering the two distinct temperatures previously mentioned. The main abundance differences are found for high G_0 PDRs, where the dust temperature varies up to ~ 30 K depending on the considered T_{dust} expression as mentioned in Section 3. In particular, these abundance differences can be of up to four orders of magnitude in the inner regions of the cloud ($A_V > 6$ mag) for the case of water and of more than six orders of magnitude for molecular oxygen. In the edge of the cloud ($A_V \leq 1$ mag), however, the abundance differences are no more than one order of magnitude.

For a high G_0 PDR with density $n = 10^5 \text{ cm}^{-3}$ (middle panels), low dust temperatures (derived from Garrod’s expression) promote the formation of OH, O_2 , and H_2O at $1 < A_V \lesssim 5.5$ mag, while for larger extinctions, high dust temperatures (obtained from Hocuk’s expression) lead to the highest oxygen-bearing molecule abundances with differences of up to seven orders of magnitude between both expressions. This is the interesting case of molecular oxygen, an elusive molecule in the ISM [Odin satellite only provided upper limits ($\leq 10^{-7}$) for its abundances especially in cold dark clouds, e.g. Pagani et al. 2003] with only a few recent detections: the massive Orion star-forming region [with $X(\text{O}_2) \sim 10^{-6}$, Goldsmith et al. 2011, Chen et al. 2014] and the low-mass dense core ρ Oph A [with $X(\text{O}_2) \sim 5 \times 10^{-8}$, Larsson et al. 2007; Liseau et al. 2012]. Recently, this molecule has also been detected in surprisingly large quantities towards the Solar system comets 67P/Churyumov–Gerasimenko (67P/C-G) by Bieler et al. (2015) with Rosetta, and in 1P/Halley by Rubin et al. (2015) with the Giotto mission. Their results confirm that O_2 is the fourth most abundant molecule in comets. In our PDR case, we find that, at $A_V \gtrsim 4$ mag, high dust temperatures allow to enhance the surface diffusion of O atoms that recombine to form solid O_2 , which is then released into the gas-phase through thermal desorption. This is in agreement with Taquet et al. (2016). We also find chemical desorption from the reaction of two solid oxygens as an important way to form O_2 gas, especially at $4 \lesssim A_V \lesssim 6$ mag (see Fig. A1, left-hand panel, in the Appendix A). We highlight the need of carrying out an O_2 search in PDRs to make quantitative comparison with our predictions.

For the particular case of water in a high G_0 PDR (middle bottom panel, Fig. 3), it presents a low abundance variation for $A_V < 1$ mag when the dust temperature varies by ~ 30 K, highlighting a gas-phase chemical formation route for this molecule via ion-chemistry at the edge of the cloud. For intermediate extinctions ($1 < A_V \lesssim 3$ mag), the H_2O abundance variations are very small (< 1 order of magnitude) between both T_{dust} expressions as also found for the OH abundances, while O_2 presents differences of about two orders of magnitude. This shows that at intermediate visual extinctions, OH is a more relevant reactant than O_2 to form water, and that the main H_2O formation route is through successive hydrogenation of atomic oxygen in agreement with Dulieu et al. (2010). In particular, we find this chemical reaction efficient for $A_V < 5$ mag (see Fig. A1, right-hand panel, in the Appendix A). For larger extinctions, the warmer the dust grains, the higher the water abundances with differences of up to four orders of magnitude between both T_{dust} , being photo and cosmic ray desorption the most efficient reactions forming gaseous water at $A_V \gtrsim 5$ mag (Fig. A1, right-hand panel, Appendix A).

If the density of the PDR increases by one order of magnitude (right-hand panels, Fig. 3), the main effect with respect to the low density case is found at the edge of the cloud (at $A_V \lesssim 0.5$ mag) where the abundances of the three molecules (OH, O_2 , and H_2O) increase by approximately two orders of magnitude for both dust temperature expressions. In the case of a low G_0 PDR (left-hand panels, Fig. 3), the temperature differences between both T_{dust} expressions are $\lesssim 10$ K. These small differences lead to variations in the abundances of OH, O_2 , and H_2O of no more than one order of magnitude for $0 \leq A_V \leq 10$ mag.

From these results, we therefore conclude that the largest impact in the chemistry of simple oxygen-bearing molecules is found in high G_0 PDRs, which present the largest dust temperature differences between the two approaches for T_{dust} . In

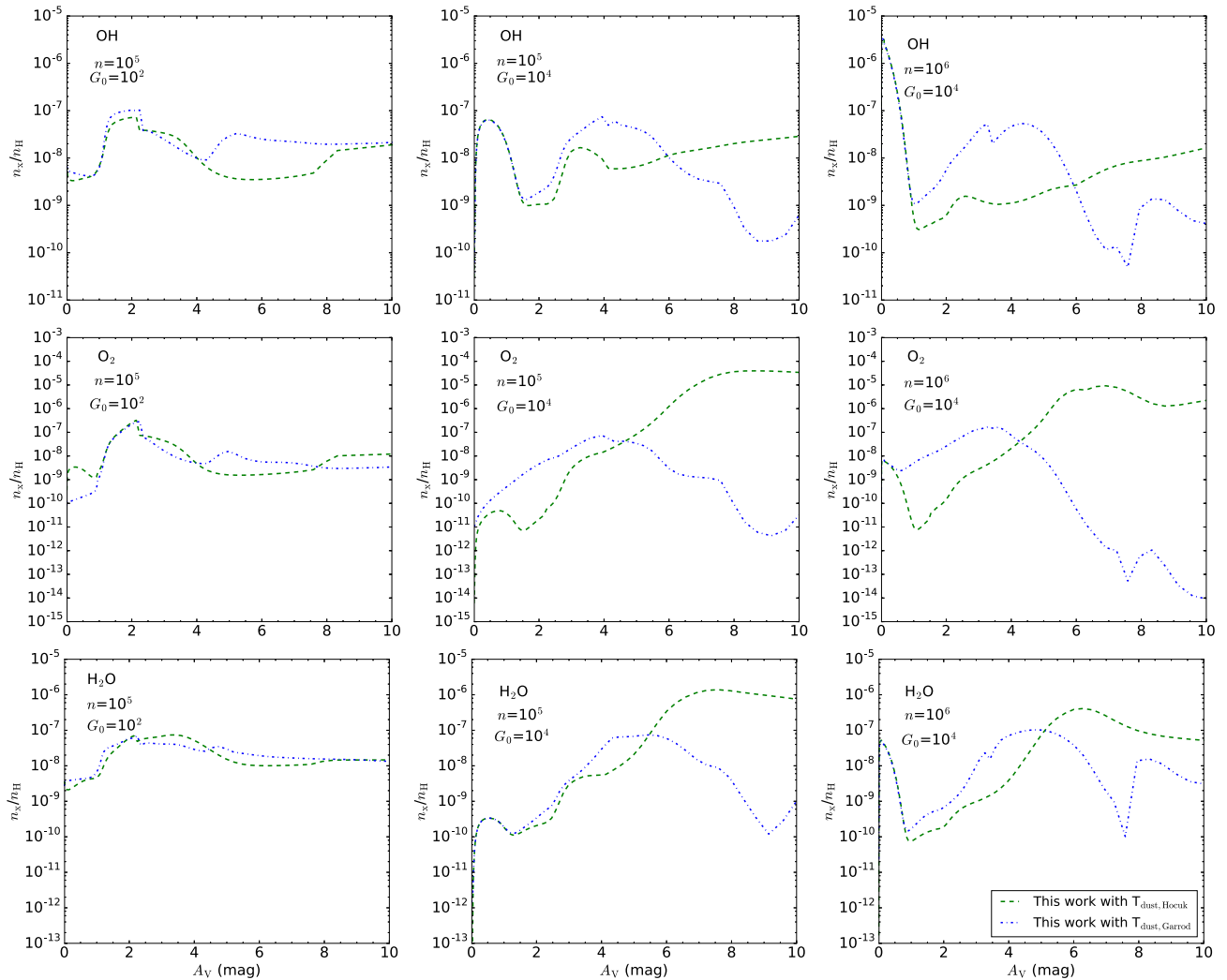


Figure 3. Abundances of OH, O₂, and H₂O obtained with the updated Meijerink PDR code using T_{dust} from Hocuk et al. (2017) (green dashed lines) and from Garrod & Pauly (2011) (blue dotted lines). Results for Model 1 are shown on the left-hand panels, for Model 2 in the middle panels, and for Model 3 on the right-hand panels.

these PDRs, low dust temperatures promote the formation of OH, O₂, and H₂O at intermediate visual extinctions ($A_V \lesssim 5$ mag), while high values of T_{dust} promote their formation at larger A_V .

5.1.2 Carbon-bearing molecules

Fig. 4 shows the abundances of carbon-bearing molecules (CH, CO, H₂CO, and CH₃OH) considering two different dust temperatures (from Hocuk’s and Garrod’s expressions). For the simplest species (CH and CO shown in the two top panels), we distinguish two regimes for any PDR type: the low visual extinction regime ($A_V \lesssim 2$ mag), where the variation of dust temperature does not have a significant impact on the abundances of these molecules since they mainly form in the gas phase, and the high visual extinction range ($A_V > 2$ mag), where their abundances can vary by up to three orders of magnitude.

For a high G_0 PDR (middle panels), Hocuk’s expression produces the highest T_{dust} values, which lead to a low CO depletion on grain surfaces and therefore to large CO gas-phase abundances (up to

two orders of magnitude larger than those obtained using Garrod’s expression). The large CO gas-phase abundance at $A_V \gtrsim 4$ mag obtained with Hocuk’s expression implies low abundances of solid CO and therefore a restriction in the formation of more complex molecules on the grain surfaces through CO ice, such as H₂CO and CH₃OH, as we observe in Fig. 4 (two bottom panels). In particular, we obtain that the abundance of H₂CO at $A_V \gtrsim 4$ mag is lower for Hocuk’s expression than for Garrod’s expression by up to approximately three orders of magnitude. This difference is even larger (up to six orders of magnitude) in the case of the complex molecule CH₃OH.

The increase of the PDR density favours the formation of all the carbon-bearing molecules at $A_V \lesssim 1$ mag as shown in Fig. 4 (right-hand panels). In particular, we find that the abundances of CH, CO, H₂CO, and CH₃OH increase by about two orders of magnitude in the edge of the cloud without finding significant differences between both T_{dust} expressions. At intermediate and large extinctions ($A_V \gtrsim 3$ mag), we observe that the density increase mainly affects the abundances obtained with the lowest T_{dust} values (blue dotted curves), with CO being the most affected molecule. In particular,

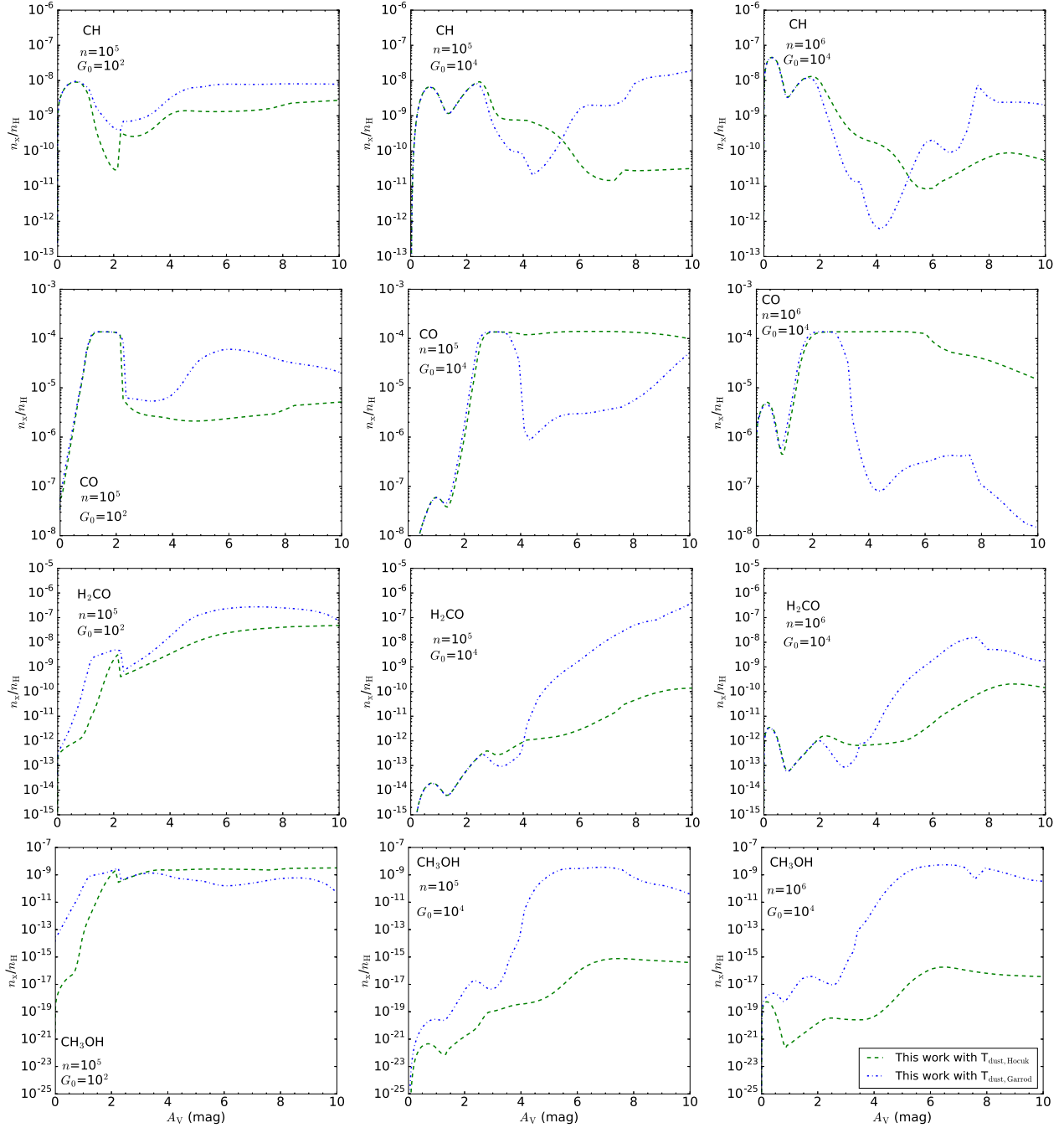


Figure 4. Abundances of CH, CO, H₂CO, and CH₃OH obtained with the updated Meijerink PDR code using T_{dust} from Hocuk et al. (2017) (green dashed lines) and from Garrod & Pauly (2011) (blue dotted lines). Results for Model 1 are shown on the left-hand panels, for Model 2 in the middle panels, and for Model 3 on the right-hand panels.

the increase of density by one order of magnitude leads to a CO abundances decrease of about three orders of magnitude due to a more efficient depletion. This promotes the formation of complex molecules. In fact, the abundances of CH₃OH are slightly larger at $A_V > 7$ mag in the PDR with density 10^6 cm^{-3} (bottom right panel) than in the PDR with $n = 10^5 \text{ cm}^{-3}$ (middle bottom panel).

For a low G_0 PDR (left-hand panels of Fig. 4), in the edge of the cloud ($A_V \lesssim 1$ mag), we only find significant differences for H₂CO and CH₃OH when changing T_{dust} , since these molecules are

mainly formed on dust grains, which makes them very sensitive to variations of dust temperature. For these two molecules, the lower the dust temperatures, the higher the abundances, since H atoms can reside on dust grains longer if temperatures are low. This is in disagreement with Le Gal et al. (2017), who suggested that the warming up of grain surfaces speeds up chemical surface processes forming complex organic molecules (COMs), explaining thus the high abundance of some COMs observed in the PDR of the Horsehead ($G_0 \sim 10^2$ and $n \sim 10^5 \text{ cm}^{-3}$, Habart et al. 2005,

Guzmán et al. 2013) with respect to the core of the cloud (Gratier et al. 2013). It must be noted that while their conclusions were deduced considering a grain warm up from ~ 10 to ~ 25 K, the T_{dust} difference in our comparison is ~ 5 K for $A_V \lesssim 1$ mag. Nevertheless, we obtain the same trend for CH_3OH in the high G_0 case (Fig. 4, middle and bottom right panels), where the T_{dust} difference for $A_V \lesssim 1$ mag is about ~ 25 K. Fig. A2 (see Appendix A) shows abundances of H_2CO and CH_3OH at $A_V \leq 1$ mag when the density is increased by one order of magnitude ($n \sim 10^6 \text{ cm}^{-3}$), while the radiation intensity remains as $G_0 \sim 10^2$. When density increases, we obtain an abundance increase for both molecules between one and three orders of magnitude. From these results, we propose an alternative stage where the presence of higher abundances of some COMs in the PDR than in the core of the Horsehead is the result of the presence of clumps with very high densities (of at least $n = 10^6 \text{ cm}^{-3}$) and low dust temperature values ($T_{\text{dust}} < 25$ K) in the edge of the cloud. See also Section 5.2.3 for a more detailed explanation of the density role in the significant enrichment of some COMs in the PDR with respect to the cloud core.

From all these results, we derive that low dust temperatures significantly promote the formation of COMs in the inner regions of high G_0 PDRs, as well as in the edge of clouds with low G_0 PDRs.

5.1.3 Solid molecules

Fig. 5 shows the abundances of solid H_2O , CO_2 , CO , CH_3OH , and CH_4 obtained using different analytical expression for T_{dust} (green dashed lines for Hocuk's expression and blue dotted lines for Garrod's expression).

For a low G_0 PDR (left-hand panels), the visual extinctions at which the first full ice monolayers of H_2O and CO_2 are formed barely changes with the T_{dust} considered due to the small difference between both expression ($\lesssim 10$ K). For both molecules, this formation occurs at $A_V \sim 2$ – 3 mag, as in diffuse molecular clouds (Boogert, Gerakines & Whittet 2015). For the case of CO , only the lowest T_{dust} leads to the formation of CO ice at intermediate and large extinctions ($A_V \geq 5$ mag), while other more complex molecules, such as CH_3OH and CH_4 , present abundances lower than 10^{-6} and do not form ice at $A_V \leq 10$ mag for any of the two T_{dust} considered. This is in disagreement with Hollenbach et al. (2009), who obtained similar maxima for the CO and CH_4 ice abundances in a PDR with $G_0 = 100$ and $n = 10^4 \text{ cm}^{-3}$, suggesting an overproduction of methane, since Öberg et al. (2008) and Boogert et al. (2015) observed solid $\text{CH}_4/\text{H}_2\text{O}$ abundances of ~ 0.05 and 0.01 in low- and high-mass young stellar objects, respectively.

In a PDR with the same density, but a G_0 two orders of magnitude higher (middle panels), the difference between both dust temperature expressions is ~ 30 K (see Fig. 1), which is high enough to make molecular depletion on to dust grains less efficient in the warmest case. This leads to the formation of H_2O and CO_2 ices at larger extinctions (between ~ 3 and 5 mag) for both T_{dust} expressions. For the case of solid CO , CH_3OH , and CH_4 , the increase of the radiation intensity from $G_0 = 10^2$ to $G_0 = 10^4$ produces a significant drop in their abundances of at least five orders of magnitude in the highest T_{dust} case (green lines), highlighting the need of cool grains to form ices of carbon monoxide, methanol, and methane.

When density increases (right-hand panels), the visual extinction at which H_2O and CO_2 ices are formed slightly decreases for both dust temperature expressions. This is due to the increase in the rate at which atoms and molecules hit dust grains, which is linearly

dependent on the gas number density. Regarding minor ice mantle components, the density increase in a very high G_0 PDR allows the formation of methane ice only at large extinctions ($A_V > 8$ mag) when the dust grain temperature remains low ($\lesssim 10$ K). No formation of methanol ice is found in any of the considered PDR types, although a low T_{dust} significantly promotes its formation.

We therefore conclude that low dust temperatures promote the formation of solid H_2O , CO , CH_3OH , and CH_4 in all type of PDRs, while warm grains promote the formation of solid CO_2 at any A_V for low G_0 , and only at very large extinctions ($A_V > 8$ mag) for high G_0 PDRs.

5.1.4 Comparison with observations: dust temperature

After the analysis of the chemical impact produced by the variation of the dust temperature using the expressions from Garrod & Pauly (2011) and Hocuk et al. (2017) in the PDR code, we compare our predictions with observations to find the temperature of the best agreement.

For the case of a low G_0 PDR, we have compared our results obtained using both T_{dust} expressions with observations of two molecules, CH_3OH and H_2CO , in the Horsehead ($n = 10^5 \text{ cm}^{-3}$ and $G_0 \sim 10^2$, Habart et al. 2005). In particular, we have compared with the ratio of these two molecules, since the estimation of their abundances with respect to H_2 presents large uncertainties due to the strong dependence of the H_2 density on the dust temperature considered.⁷ Fig. 6 shows this comparison considering observations of the H_2CO and CH_3OH in the PDR (the IR peak at $A_V \sim 1$ mag) and the core ($A_V \sim 8$ mag, Pety et al. 2012). The kinetic temperatures assumed to infer the observational results were $T_{\text{kin}} = 40$ – 65 and 20 K, for the PDR and the core respectively (Guzmán et al. 2011, 2013), which are consistent with the PDR model temperatures for both regions (Fig. 1, right-hand panel). The results show that the $\text{CH}_3\text{OH}/\text{H}_2\text{CO}$ ratio in the PDR region is reproduced by either expressions, however none of them reproduces the observations in the core. Nevertheless, the difference between observations and model is about one order of magnitude using Hocuk's expression, and about three orders of magnitude using Garrod's expression at $A_V = 8$ mag.

For the case of a high G_0 PDR, observations of the densest parts of the Orion Bar ($n = 10^5$ – 10^6 cm^{-3} and $G_0 \sim 10^4$, Marconi et al. 1998, Leurini et al. 2010) carried out with the Herschel space telescope reveal a dust temperature gradient from ~ 70 to ~ 48 K for the largest grains at different positions in the Bar (Arab et al. 2012). Millar & Williams (1993) also show through FIR observations that the temperatures of dust grains with size ~ 3000 Å in the Bar region are about 75 K. Comparing these results with those shown in Fig. 1 (left-hand panel), we clearly see that the T_{dust} expression from Hocuk et al. (2017) provides dust temperature values in full agreement with the observations of the Orion Bar. In the following, we consider the dust temperature expression from Hocuk et al. (2017).

5.2 Time-dependent effects

Figs 7–10 show the abundances of several families of molecules as a function of time ($10^4 \leq t \leq 10^7$ yr) and visual extinction ($0 \leq A_V \leq 10$

⁷ Leurini et al. (2010) found a variation in the density of H_2 larger than a factor of 2 when the difference considered in the dust temperature is 20 K.

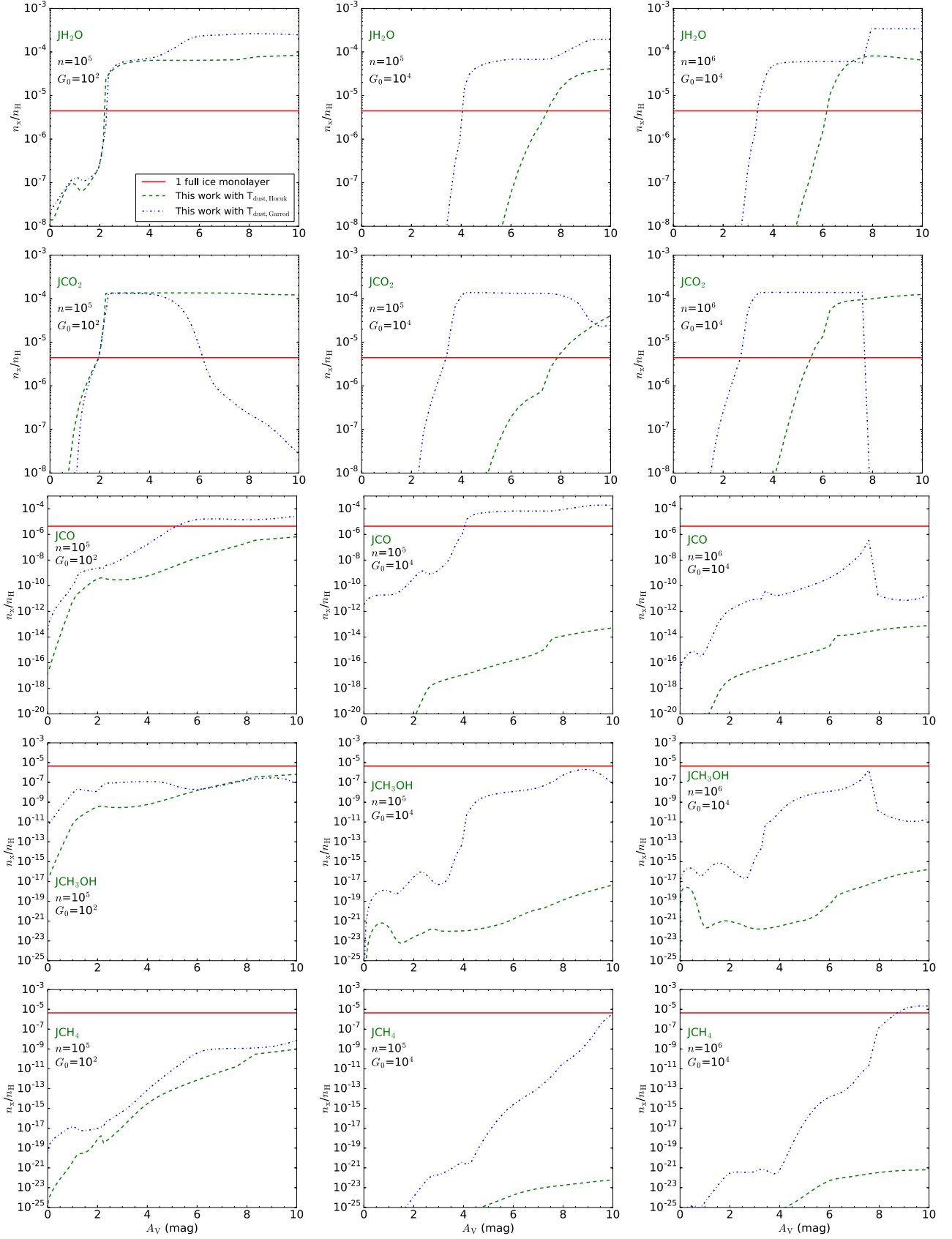


Figure 5. Abundances of JH₂O, JCO₂, JCO, JCH₃OH, and JCH₄ obtained with the updated Meijerink PDR code using T_{dust} from Hocuk et al. (2017) (green dashed lines) and from Garrod & Pauly (2011) (blue dotted lines). Results for Model 1 are on the left-hand panels, for Model 2 in the middle panels, and for Model 3 on the right-hand panels. J_i means solid i . The red solid line represents the number of possible adsorption sites on grain surfaces per cm².

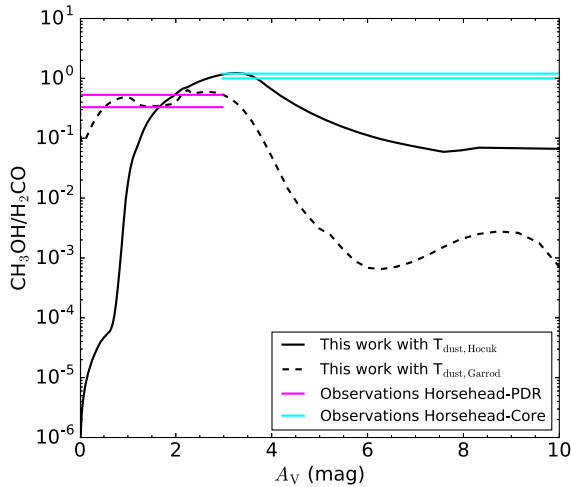


Figure 6. $\text{CH}_3\text{OH}/\text{H}_2\text{CO}$ ratio obtained for a PDR with $G_0 = 10^2$ and $n = 10^5 \text{ cm}^{-3}$ considering two different T_{dust} expression: from Garrod & Pauly (2011) (black dashed line) and from Hocuk et al. (2017) (black solid line). Observations (Guzmán et al. 2011, 2013) of the PDR and the core of the Horsehead are also shown with magenta and cyan lines, respectively, considering their uncertainties through a double line.

mag) for three different type of PDRs (Models 1, 2, and 3 defined in Section 5).

5.2.1 Simple oxygen-bearing molecules

Fig. 7 shows the time evolution of the chemical abundances of OH (top), O_2 (middle), and H_2O (bottom). For any type of PDR, we obtain that the abundances of these three molecules at the edge of the cloud are $\lesssim 10^{-8}$ for any evolutionary time. However, as the visual extinction increases, their abundances increase with a difference of up to 10 orders of magnitude between the edge ($A_V \leq 1 \text{ mag}$) and the inner ($A_V > 6 \text{ mag}$) part of the cloud depending on the type of PDR.

For a low G_0 PDR (left-hand panel), although these three molecules present their highest abundances at $A_V \gtrsim 6 \text{ mag}$, there are significant time differences between them. In the case of water, the abundance peak ($\sim 10^{-5}$) is reached at an early evolutionary stage ($t \sim 10^4 \text{ yr}$), indicating that the reactions forming gas-phase water are fast. This abundance peak only presents variations lower than one order of magnitude for $A_V \gtrsim 6 \text{ mag}$ until $t \sim 10^6 \text{ yr}$, while for longer times water gas starts being significantly destroyed to form water ice. This abundance decrease is also found for O_2 and OH, which, after depletion, represents an important reactant to form CO_2 ice through the surface reaction $\text{JOH} + \text{JCO} \rightarrow \text{JCO}_2 + \text{JH}$. This is in agreement with Hollenbach et al. (2009) who also found that most of the gas-phase oxygen goes to H_2O ice and CO_2 ice at $t \sim 10^7 \text{ yr}$ for $A_V > 8 \text{ mag}$. Long time-scales promote therefore the destruction of simple oxygen-bearing molecules at intermediate and large extinctions.

When the intensity of the radiation field increases by two orders of magnitude (middle panels, Fig. 7), the time for which the maximum abundances of H_2O are reached also increases by about one order of magnitude ($t \sim 10^5 \text{ yr}$). Similar behaviour is found for the other two species (OH and O_2), which indicates that, in high G_0 PDR, long time-scales promote the formation of simple oxygen-bearing molecules only in the inner regions of the cloud. In this case, the maximum abundances found for H_2O and OH are about one order

of magnitude lower than those found in the low G_0 PDR case. Molecular oxygen, however, reaches the same maximum abundance ($\sim 10^{-5}$) independently on the intensity of the radiation, time being the main difference at which this value is reached. This shows that water formation is more linked to successive hydrogenations of atomic oxygen than to reactions involving molecular oxygen, as also found in Section 5.1.1.

The main effect of increasing the density by one order of magnitude in a high G_0 PDR is found for H_2O (bottom right panel, Fig. 7). For this molecule, this increase allows water to reach its maximum abundance at an early evolutionary stage ($t < 10^4 \text{ yr}$), while in the lower density PDR (middle bottom panel) its maximum abundance is reached at $t \gtrsim 10^5 \text{ yr}$. At low visual extinctions ($A_V < 5 \text{ mag}$), the abundances of these three simple oxygen-bearing molecules remain very low ($\lesssim 10^{-8}$) at any evolutionary stage.

From these results, we deduce that at intermediate and large A_V , long time-scales promote the formation of simple oxygen-bearing molecules in high G_0 PDRs, while favour their destruction if G_0 is low. In the edge of the cloud, no significant time effects are found on the evolution of these species.

5.2.2 Simple carbon-bearing molecules

Fig. 8 shows the chemical evolution of CH (top), CO (middle), and H_2CO (bottom) as a function of time and visual extinction. We obtain that CH is mainly formed at the edge of the cloud for either a low and a high G_0 PDR (top left and top middle panels), while in the core its presence is much less significant. For a high intensity radiation field (top middle panel), abundances of CH barely change with time for $A_V \lesssim 2 \text{ mag}$. In this case, CH mainly forms through the very endothermic reaction between H_2 and the ion C^+ , which is efficiently formed with high radiation intensity. When G_0 decreases (top left-hand panel), the abundance of C^+ also decreases. In this case, we find that CH abundances progressively increase with time.

Unlike CH, we do not find significant time dependence in the abundances of CO at the edge of the three considered PDRs. For this molecule, time only becomes important at $A_V \gtrsim 3 \text{ mag}$ and presents different effects depending on the type of PDR. For a low G_0 PDR (left middle panel, Fig. 8), long time-scales promote its destruction since it freezes out, while in a high G_0 PDR (middle panel) the opposite behaviour is found. We also obtain that formaldehyde presents a similar time dependence (see left and middle bottom panels). In the case of CO, the abundance variation due to time effects is only $\lesssim 1$ order of magnitude for any A_V , suggesting that this species is very stable with time. In other words, for each visual extinction, it is formed almost as much CO as is destroyed for any type of PDR. Similar results are also found for H_2CO , whose abundances change by no more than two orders of magnitude over 10^7 yr , especially at low and intermediate A_V .

When the density of the PDR increases by one order of magnitude (right-hand panels), we can distinguish two regimes. For a low extinction regime ($A_V \lesssim 1 \text{ mag}$), the main effect found is the increase of the abundances of the three species (CH, CO, and H_2CO) by up to two orders of magnitude, since, close to the edge, destruction of molecules is dominated by photodissociation whose rates varies as n , while formation rates vary as n^2 , resulting in abundances that increase with density. In a higher extinction regime, the main effect is found on the abundances of CO at $A_V > 5 \text{ mag}$. In particular, while CO reaches its maximum abundance at $t \gtrsim 10^5 \text{ yr}$ in the lowest density PDR (middle panel), this value is

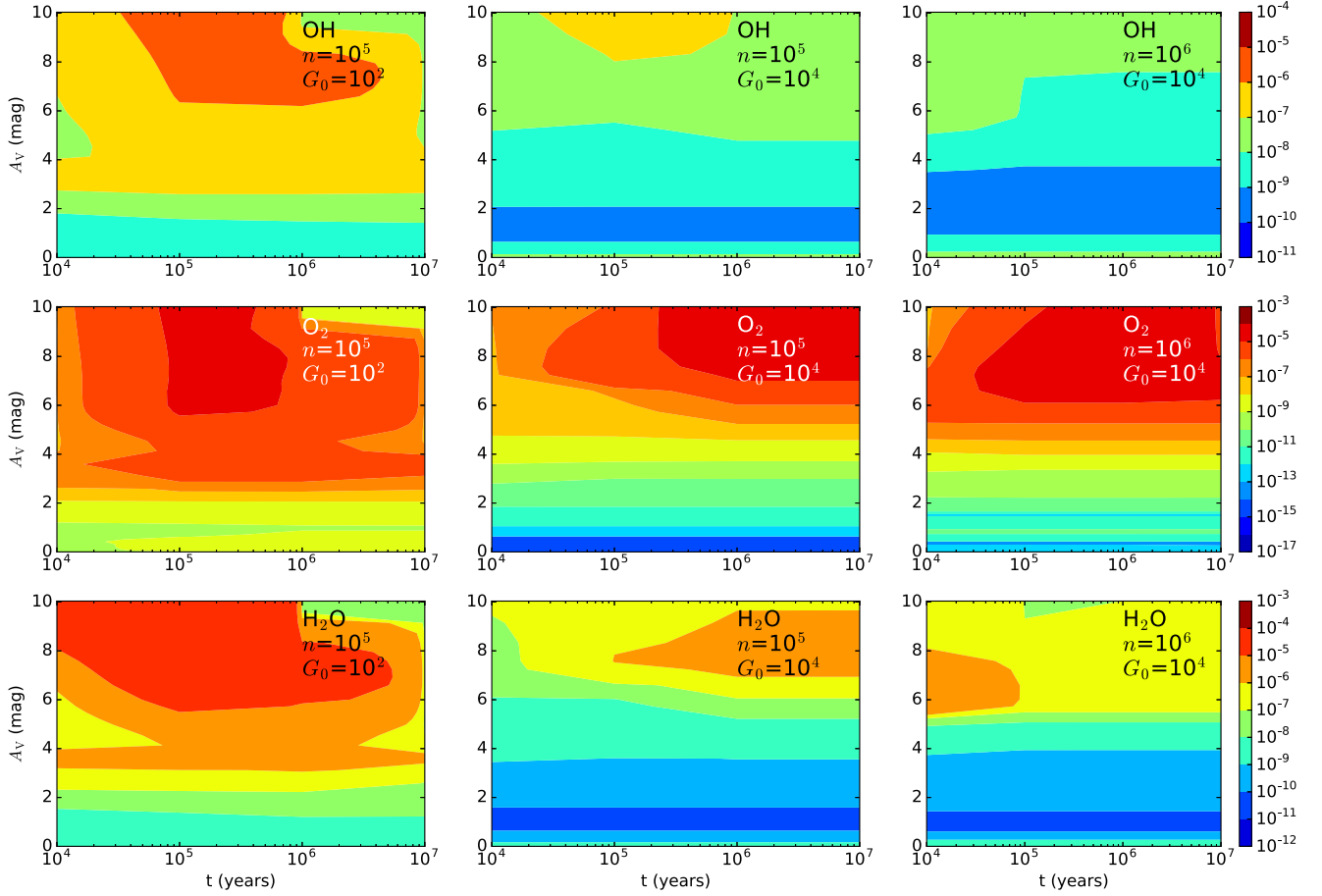


Figure 7. Contour maps with the abundances of OH, O₂, and H₂O with respect to H nuclei for Models 1 (left-hand panel), 2 (middle panel), and 3 (right-hand panel) as a function of time (*x*-axis) and visual extinction (*y*-axis).

reached at $t < 10^5$ yr in the high-density PDR. This is due to a variation in the efficiency of the chemical reactions forming CO. In the low density case, CO mainly forms through dissociative recombination of HCO^+ , which is formed through the reaction between C^+ and H_2O (H_2O being more efficiently formed at large extinctions and long time-scales as shown in Fig. 7, middle bottom panel). However, as the density increases, the formation of HCO^+ through the ionization of chemically desorbed HCO becomes more efficient.

5.2.3 Complex organic molecules

Fig. 9 shows the chemical evolution of CH_3OH , CH_3CN , and CH_2CO as a function of time and visual extinction.

The CH_3OH formation starts being efficient at $A_V > 2$ mag, since it mainly forms on grain surfaces through chemical desorption upon the surface reaction between solid H with solid H_3CO . In a low G_0 PDR, the maximum abundances of methanol are obtained at $10^5 \lesssim t \lesssim 5 \times 10^6$ yr, when enough time has passed to form solid H_3CO , its precursor. In the case of considering a more intense radiation field (two orders of magnitude higher, top middle panel), we obtain that the abundances of methanol sharply decrease at any visual extinction and evolutionary stage by several (up to 10) orders of magnitude. Only the increase of density (top right-hand panel, Fig. 9) promotes the formation of methanol deep in the cloud at $t < 10^5$ yr.

Results for molecule CH_3CN are shown in the middle panels of Fig. 9. In the gas phase, one of the principal precursors of CH_3CN is HCN, which mainly forms through an exchange chemical reaction whose activation energy barrier is 100 K (MacKay 1999). This barrier is slightly higher than the gas temperature of the low G_0 PDR at low extinctions (see right-hand panel of Fig. 1), which explains the low ($\lesssim 10^{-16}$) abundances of methyl cyanide at the edge of this type of PDRs (left middle panel in Fig. 9). As A_V increases, the abundance of CH_3CN increases by up to 10 orders of magnitude in the core, indicating that this molecule is significantly enhanced by reactions occurring on grain surfaces. Regarding time effects, Fig. 9 (left-hand panel) shows that CH_3CN presents differences in its abundances no larger than one order of magnitude over time at $A_V \gtrsim 3$ mag. The opposite effect is, however, found when G_0 increases by two orders of magnitude (middle panel). In this case, time effects become more important for the evolution of CH_3CN as the visual extinction increases. In particular, its abundances increase by up to three orders of magnitude from $t = 10^4$ to $t = 10^6$ yr, showing that CH_3CN is a late-forming molecule in high G_0 PDRs.

Particularly interesting is the effect of increasing the density of the PDR (right middle panel, Fig. 9) by one order of magnitude for CH_3CN at $A_V \lesssim 0.5$ mag. This produces an increase of the CH_3CN abundances of at least two orders of magnitude with respect to the low density case (middle panel), becoming even higher than some values found at intermediate extinctions ($A_V \sim 5$ mag) for an early ($t < 10^5$ yr) evolutionary stage. We find that this effect, which

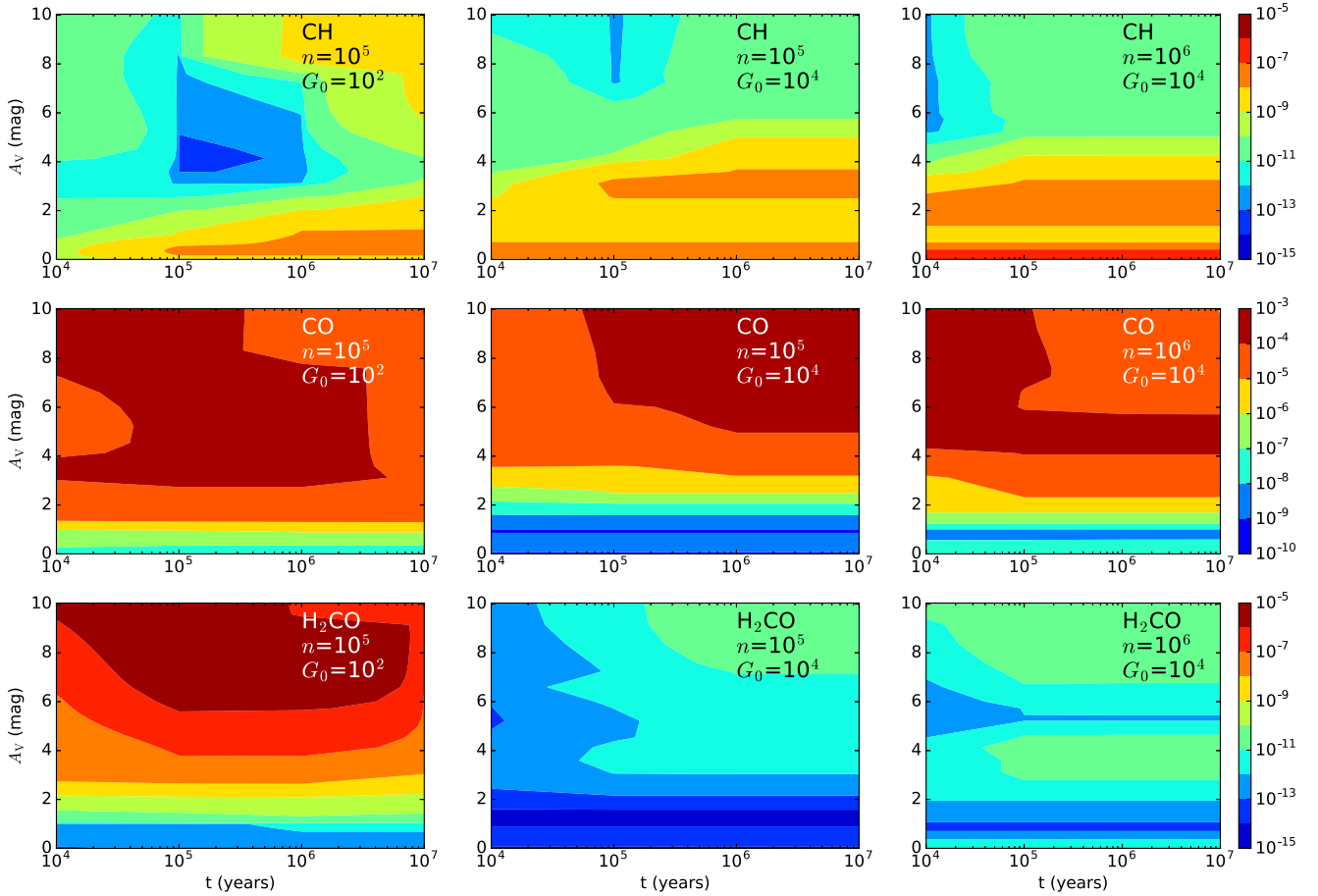


Figure 8. Contour maps with the abundances of CH, CO, H₂CO, CH₃OH, and CH₃OH with respect to H nuclei for Models 1 (left-hand panel), 2 (middle panel), and 3 (right-hand panel) as a function of time (x-axis) and visual extinction (y-axis).

has also been observationally detected by Gratier et al. (2013), is mainly due to a more efficient formation of HCN (precursor of CH₃CN in the gas phase) in the edge of the cloud produced by the density increase. In particular, at $A_V \lesssim 0.5$ mag, we obtain an increase of HCN of about two orders of magnitude when the density increases by one order of magnitude (comparison between middle and right-hand panels of Fig. A3 in the Appendix A). In this case of a very high-density PDR, CH₃CN mainly forms through radiative association ($\text{CH}_3^+ + \text{HCN}$, with CH_3^+ being quite abundant at low A_V with respect to large extinctions due to the high UV radiation), followed by dissociative recombination.

Ketene has long been identified in the ISM and different gas-phase pathways (with ethylene ions as precursors) have been proposed for its formation (Millar, Herbst & Charnley 1991). However, its detection in the cold pre-stellar cores L1544 (Spezzano et al. 2017) and L1689B at temperatures of ~ 10 K suggests a formation with grain surface chemistry (through methane–carbon monoxide ices) and subsequent non-thermal desorption via induced UV photons and cosmic ray impacts (Bacmann et al. 2012, Maity, Kaiser & Jones 2014). In a low G_0 PDR (bottom left panel, Fig. 9), we find the maximum CH₂CO abundances at late times ($5 \times 10^5 \lesssim t \lesssim 5 \times 10^6$ yr) in the inner regions ($A_V > 6$ mag) of the PDR.

As G_0 increases (middle bottom panel), the abundance of CH₂CO sharply decreases for all visual extinctions and for any evolutionary stage due to the increase of radiation, which prevents the formation

of ketene ice precursors. In this case, the formation of ketene at the edge of the PDR becomes inefficient and time effects are only important at intermediate and large extinctions. In general, we observe that abundances of CH₂CO change by no more than one order of magnitude at $t > 10^5$ yr for each visual extinction, independently on the type of PDR. We therefore deduce that visual extinction is a more important factor than time for the formation of ketene.

5.2.4 Solid molecules

The two top rows of Fig. 10 show the chemical evolution of solid water (JH₂O) and solid carbon dioxide (JCO₂) for Models 1, 2, and 3, as a function of time.

For a low G_0 PDR (top left panel), the abundances of solid water increase with visual extinction. In particular, the maximum abundance ($\sim 10^{-4}$) of JH₂O with respect to hydrogen is reached at $t \gtrsim 10^5$ yr for $A_V \sim 5$ –8 mag and remains roughly constant over more than one million years. When G_0 increases, radiation effects prevent solid water formation at low A_V . In this case, the maximum solid water abundance is roughly the same as in the low G_0 PDR, but found at larger extinctions ($A_V > 7$ mag). We also find that this abundance peak is reached at an earlier evolutionary stage ($\sim 5 \times 10^4$ yr) than in the low G_0 case. This time becomes significantly lower as the density of the PDR increases (right top panel). We therefore

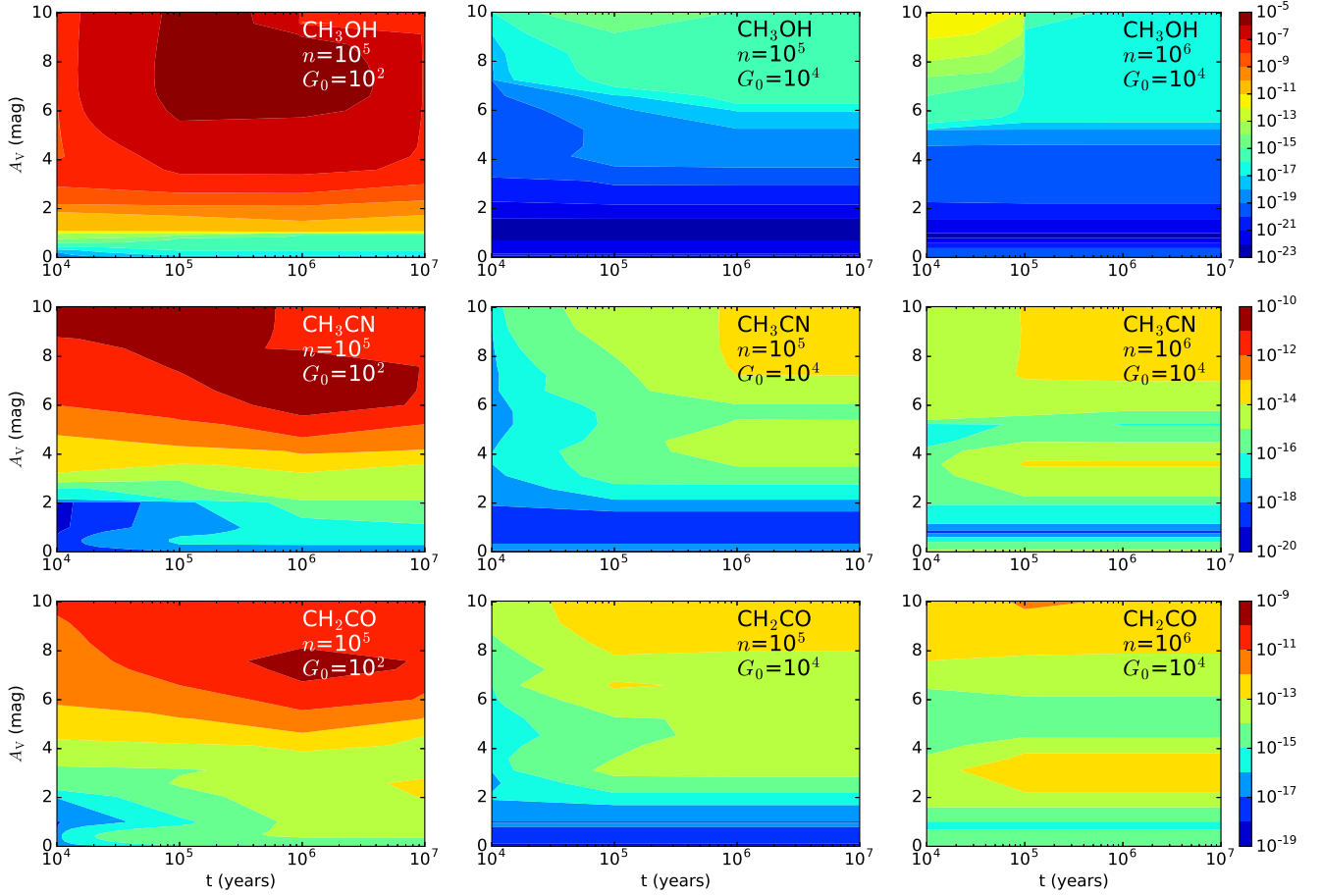


Figure 9. Contour maps with the abundances of CH_3OH , CH_3CN , and CH_2CO with respect to H nuclei for Models 1 (left-hand panel), 2 (middle panel), and 3 (right-hand panel) as a function of time (x-axis) and visual extinction (y-axis).

conclude that high G_0 values promote the formation of solid water at earlier evolutionary stages and larger extinctions than low G_0 values.

Fig. A4 (Appendix A) shows the threshold to form one full monolayer of water ice (see Esplugues et al. 2016 for more details about the calculation of this limit), together with the fractional water abundances over time. The first water ice monolayer is formed at a very early stage (10^4 yr) for any type of PDR, although the visual extinction varies between 3 and 4 mag depending on the considered G_0 value (the higher G_0 , the larger A_V due to the increase of UV radiation, which prevents ice formation). For longer time-scales, we also find formation of full monolayers of water ice for any PDR, but at larger extinctions than in the early stage.

Results for solid CO_2 are also shown in Fig. 10 (second row). The abundances of this species for a low G_0 PDR (left-hand panel) at $A_V < 1$ mag are very low ($\lesssim 10^{-10}$) and independent on the evolutionary stage. By contrast, for larger extinctions, abundances of JCO_2 present a strong time dependence with variations of up to four orders of magnitude between the early and the evolved stages. The maximum abundance ($\sim 10^{-5}$) of solid CO_2 is first reached very deep in the cloud ($A_V > 8$ mag) at $10^5 < t < 10^6$ yr, while for a more evolved cloud ($10^6 \lesssim t \lesssim 10^7$ yr), this abundance peak is reached at much lower extinctions ($A_V \lesssim 3$ mag). In the late stage, we find that the formation of JCO_2 mainly

occurs through the reaction between solid CO and solid O at low A_V , but, as the visual extinction increases, we obtain that solid OH also becomes an important precursor of JCO_2 . For this type of low G_0 PDR, the maximum number of full CO_2 ice monolayers is reached at a late stage ($t = 10^6$ yr), according to results from Fig. A5 (Appendix A).

For a higher G_0 PDR (Fig. 10, second row, middle panel), we obtain that the abundances of solid CO_2 are only significant at $A_V \gtrsim 5$ mag and that they increase as the cloud evolves. We also find that the density increase promotes the formation of JCO_2 at earlier stages (bottom right panel) than the low density case, as also found for JH_2O , allowing to reach JCO_2 abundances $\gtrsim 10^{-5}$ at $t \sim 5 \times 10^4$ yr at intermediate extinctions.

Comparing the results for both molecules (solid H_2O and solid CO_2), we deduce that carbon dioxide is a more time-dependent species than water in low G_0 PDRs. We also derive that CO_2 is a late-forming ice with respect to water ice in dense PDRs, since the formation of the first water ice monolayer occurs at $t \sim 10^4$ yr, while for CO_2 occurs at $t \sim 10^5$ – 10^6 yr for $A_V \leq 10$ mag. We here highlight that ices form in layers, with water ice as first layers, and CO_2 ice on the top in PDRs.

Other ices, such as CO and CH_3OH (Fig. 10, the two bottom rows), present significant abundances only in the low G_0 case at large visual extinctions ($A_V > 6$ mag) for $t \gtrsim 10^6$ yr. Nevertheless, their abundance peaks are $\sim 10^{-6}$ – 10^{-8} , i.e. up to four orders

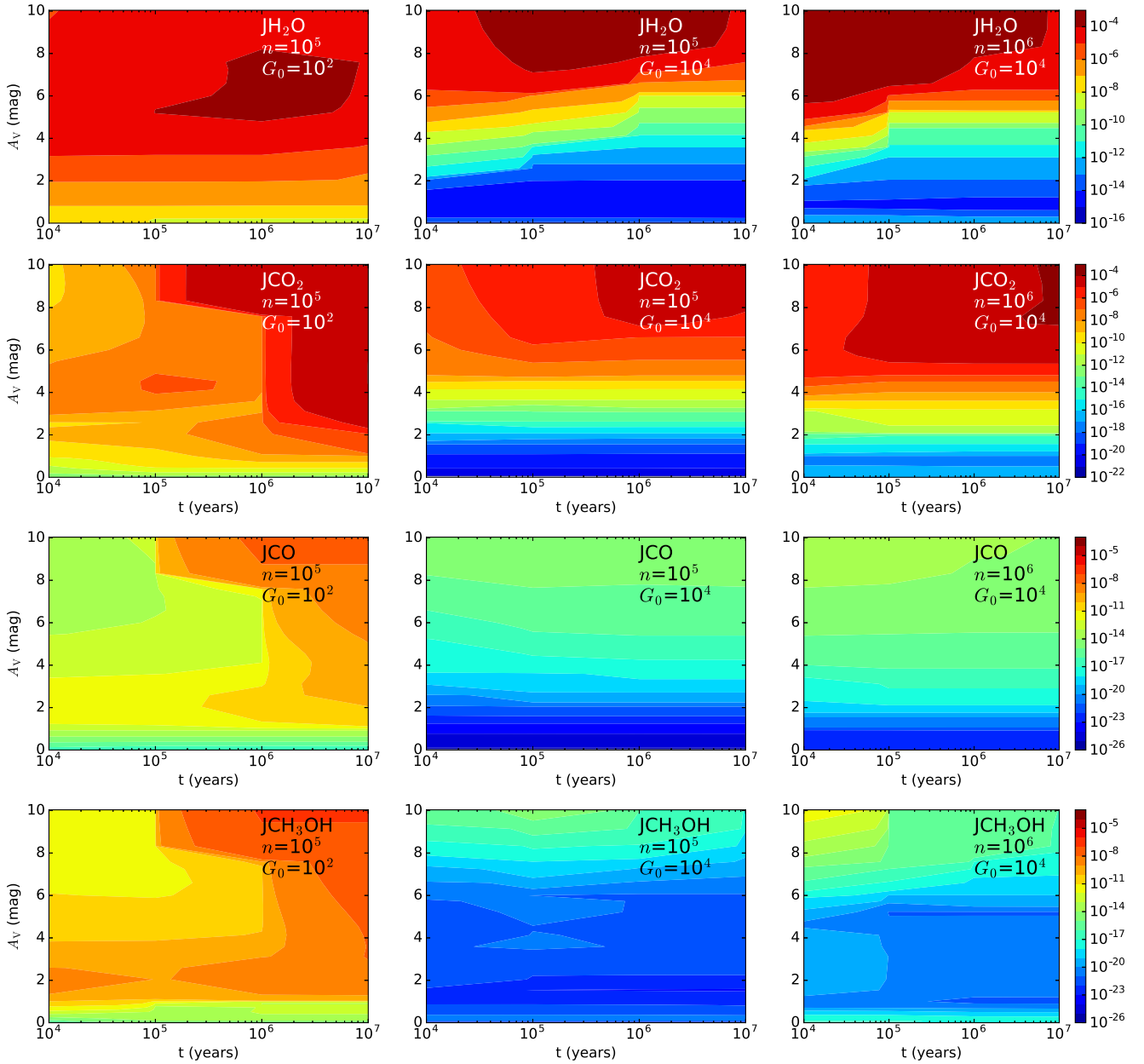


Figure 10. Contour maps with the abundances of solid H_2O , CO_2 , CO , and CH_3OH with respect to H nuclei for Models 1 (left-hand panel), 2 (middle panel), and 3 (right-hand panel) as a function of time (x-axis) and visual extinction (y-axis).

of magnitude lower than the maximum water ice abundance. They are therefore minor and late-forming ice constituents in PDRs.

5.2.5 Steady state

A chemical system reaches equilibrium when the rate at which each molecule is formed is equal to the rate at which it is destroyed, keeping its abundance constant over time. As previously stated, at low visual extinctions ($A_V \lesssim 1$ mag) in a molecular cloud, the energy balance is dominated by FUV photons and the chemical time-scales are very short compared to the molecular cloud lifetime (10^6 – 10^7 yr). However, as the visual extinction increases, certain

chemical time-scales become comparable to cloud lifetimes and steady-state chemistry does not apply. Time at which steady state is reached can also be affected by several mechanisms, such as turbulent motions (which can mix external regions exposed to the UV field with the inner regions of the cloud), star formation and the violent phenomena associated to its early stages. Here we only analyse when steady state is reached depending on the PDR type.

In Figs 7–10, we have shown the chemical evolution of different molecule families over time (10^4 – 10^7 yr) for low and high G_0 PDRs, and also varying the density. For the low visual extinction case ($A_V \lesssim 1$ mag), we observe that steady state is reached at early times ($t \lesssim 10^5$ yr) by all the considered molecules in the different PDRs, and, in particular, at $t < 10^4$ yr in high G_0 PDRs. Only

Table 3. Observational abundances with respect to total hydrogen nuclei in the Horsehead and the Orion Bar.

Species	Horsehead	Core	Orion Bar
	PDR		
H ₂ CO	$(2.9 \pm 0.4) \times 10^{-10}$	$(2.0 \pm 0.3) \times 10^{-10}$	$(1.8 \pm 0.9) \times 10^{-9}$
CH ₃ OH	$(1.2 \pm 0.2) \times 10^{-10}$	$(2.3 \pm 0.3) \times 10^{-10}$	$(1.5 \pm 0.9) \times 10^{-9}$
CO	$(5 \pm 3) \times 10^{-5}$	–	$(1.5 \pm 0.6) \times 10^{-4}$
CH	–	–	$(6.0 \pm 0.9) \times 10^{-8}$
H ₂ O	–	–	$(9 \pm 3) \times 10^{-10}$

Data for the Horsehead are obtained from Pety et al. (2005) and Guzmán et al. (2013, 2014). Data for the Orion Bar are obtained from Nagy et al. (2017) and Cuadrado et al. (2017). Abundances for CO, CH, and H₂O have been obtained considering $N(\text{H}) = 3 \times 10^{21} \text{ cm}^{-2}$ (van der Werf, Goss & O'Dell 2013).

a few complex molecules (CH₃CN and CH₂CO) present a slower chemical evolution in the $G_0 = 100$ case with equilibrium times $10^5 \lesssim t < 10^6$ yr.

For larger extinctions, however, chemical equilibrium is reached at very different times, which strongly depends on the PDR characteristics. In a high G_0 PDR with density $n = 10^5 \text{ cm}^{-3}$ (middle panels of Figs 7–10), steady state is reached at $t \lesssim 10^6$ yr for all molecules in the range $0 \leq A_V \leq 10$ mag. If the density is increased by one order of magnitude (right-hand panels), the chemical equilibrium is reached even at shorter times ($t < 5 \times 10^5$ yr) for most of the species. By contrast, in a cloud associated to a PDR with low intensity radiation field (left-hand panels, Figs 7–9), chemical equilibrium is reached at very long time-scales ($10^6 \lesssim t \lesssim 10^7$ yr) for $A_V > 2$ mag, i.e. at a time comparable to the cloud lifetime. This large time difference to reach equilibrium is mainly due to the temperature variation between both (low and high G_0) PDRs; in the low G_0 case, the temperature is significantly lower than in the high G_0 case (by ~ 50 and ~ 25 K for the gas temperature at 3 and 8 mag, respectively, and by ~ 25 and ~ 20 K for the dust temperature at those A_V , Fig. 1), producing a decrease in the reaction rate, and some chemical barriers cannot be overcome.

5.2.6 Comparison with observations: abundances

In this section, we compare our model abundances with observations (Table 3) of CO, CH, H₂O, H₂CO, and CH₃OH in the Horsehead ($G_0 \sim 100$ and $n \sim 10^5 \text{ cm}^{-3}$, Habart et al. 2005, Guzmán et al. 2013) and the Orion Bar ($G_0 \sim 10^4$ and $n \sim 10^4$ – 10^6 cm^{-3} , Marconi et al. 1998, Leurini et al. 2010). According to these physical conditions, our Model 1 would correspond to the Horsehead PDR and Model 3 to the Orion Bar.

For the case of simple molecules in the Horsehead PDR, Pety et al. (2005) derived a C¹⁸O abundance of 1.9×10^{-7} in the IR peak ($A_V \sim 1$ mag). Tercero et al. (2010) and Esplagues et al. (2013) obtained a ¹⁶O/¹⁸O ratio ~ 250 , which is lower than the Solar isotopic abundance (~ 500 , Anders & Grevesse 1989). Considering a ¹⁶O/¹⁸O ratio of 250, we derive an abundance for CO in the Horsehead PDR of 5×10^{-5} . We reproduce this value in the Model 1 at a visual extinction of $1 \lesssim A_V < 3.5$ mag for any evolutionary time (Fig. 8, left middle panel), in agreement with results from Pety et al. (2005). In the Orion Bar, Nagy et al. (2017) observed CO, CH, and H₂O with abundances of 1.5×10^{-4} , 6×10^{-8} , and 9×10^{-10} , respectively. We reproduce these values with the Model 3 ($n = 10^6 \text{ cm}^{-3}$ and $G_0 = 10^4$) at $A_V \lesssim 2.5$ mag for any evolutionary stage as well, since the abundances of these species barely change over time in this PDR model. The T_{kin} considered in Nagy et al. (2017) to obtain the observational CO abundance is ~ 137 K, in agreement

with the gas temperature considered in our model (Fig. 1, right-hand panel, solid black line) for the range ($A_V \lesssim 2.5$ mag) where the observations are reproduced.

The H₂CO abundance (2.9×10^{-10}) in the Horsehead PDR was observed at the IR peak at $A_V \sim 1$ mag (Guzmán et al. 2011, Pety et al. 2012, Guzmán et al. 2013). We reproduce this value for any evolution time at $A_V \sim 1.5$ mag (see Fig. 8, bottom left panel), which represents an extinction of about 1 mag lower than in Esplagues et al. (2016). The H₂CO observational abundance was derived using a non-local excitation and radiative transfer model considering $n_{\text{H}} = 5 \times 10^4$ – 10^5 cm^{-3} , assuming a kinetic temperature $T_{\text{kin}} = 40$ – 65 K, which are consistent with our PDR model parameters (Fig. 1, right-hand panel).

The observed H₂CO abundance in the core ($A_V \sim 8$ mag, Pety et al. 2012, Guzmán et al. 2013) of the Horsehead is $\sim 2 \times 10^{-10}$ (obtained considering $T_{\text{kin}} = 20$ K), however we predict abundances at least three orders of magnitude higher than this value at $3 < A_V \leq 10$ mag. The dust temperature considered in our PDR model (slightly lower than 20 K, Fig. 1 left-hand panel) could be a main factor producing this overestimation, since the typical dust temperature considered in this extinction range of the Horsehead is $T_{\text{dust}} \sim 20$ – 30 K (Goicoechea, Compiègne & Habart 2009, Guzmán et al. 2013), and the lower the dust temperature, the larger the H₂CO abundances at intermediate and large extinctions as found in Section 5.1.2. In particular, we find that a T_{dust} difference of only ~ 4 K (difference found at $A_V = 8$ mag from Fig. 1) leads to an H₂CO abundance difference of one order of magnitude (Fig. 4).

For the Orion Bar case, Leurini et al. (2006, 2010) observationally deduced that H₂CO traces the warm interclump close to the strong FUV-field in the Orion Bar. We obtain a difference between the observed and the modelled H₂CO abundance of less than two orders of magnitude at $A_V \sim 2.5$ – 4.5 mag and $t \geq 5 \times 10^4$ yr, for a high (10^6 cm^{-3}) density PDR model (see Fig. 8, bottom right panel). This difference is much lower than that obtained in Esplagues et al. (2016) and similar to that obtained by Cuadrado et al. (2017), who also found that H₂CO survives in the extended gas directly exposed to the strong FUV flux. Observational H₂CO in the Orion Bar was derived using a non-LTE LVG model with $T_{\text{kin}} = 150$ – 250 K, $T_{\text{dust}} \geq 60$ K, and $n(\text{H}_2) = 10^6 \text{ cm}^{-3}$ (Cuadrado et al. 2017), consistent with our density, dust and gas model temperatures.

The observed abundance of methanol in the PDR of the Horsehead (at $A_V \sim 1$ mag) is 1.2×10^{-10} (Guzmán et al. 2013). We reproduce this value at $A_V \sim 1.5$ – 2.5 mag independently on the stage of evolution, since our models show that CH₃OH is formed as fast as is destroyed for this visual extinction range over time (see Fig. 9, top left panel). In the case of the cloud core of the Horsehead, however, we obtain an overestimation of the CH₃OH abundance with respect to the observed value (2.3×10^{-10} at $A_V \sim 8$ mag) of at least two orders of magnitude.

In the case of the Orion Bar, Leurini et al. (2006, 2010) deduced that CH₃OH traces the denser and cooler clumps observed in its inner region. The observed CH₃OH abundance in the Orion Bar is 1.5×10^{-9} (Table 3). As previous studies (e.g. Cuadrado et al. 2017), we also underestimate this value by several orders of magnitude (see Fig. 9, right top panel). At present, no model seems to reproduce the inferred abundances of CH₃OH or H₂CO towards the Orion Bar. Nevertheless, it is interesting to highlight the effect of increasing by one order of magnitude the density of a high G_0 PDR model. It leads, in an early stage ($t < 10^5$ yr), to a sharp increase of the CH₃OH abundance of about six orders of magnitude between $A_V = 4$ and $A_V = 9$ mag, suggesting that the

observations of methanol in the Orion Bar would correspond to the presence of a very dense clump ($n \geq 10^7 \text{ cm}^{-3}$) formed at an early stage ($t \lesssim 10^5 \text{ yr}$). On the other hand, turbulent diffusion could also be an important process, directly affecting the abundances of methanol and leading to these differences between model results and observations, since this mechanism significantly increases the abundances of its precursors (CO) in the inner parts of the cloud (Bell et al. 2010).

6 SUMMARY AND CONCLUSIONS

We have presented for the first time a study about the effects of dust temperature and time on the chemistry of several types of PDRs using an updated version of the Meijerink PDR code.

Considering two distinct dust temperature expressions (from Garrod & Pauly 2011, and Hocuk et al. 2017), which differ from each other up to $\sim 30 \text{ K}$ depending on the PDR characteristics, we have found the most significant chemical impact in high G_0 PDRs. In this case, the formation of complex molecules, such as methanol, is clearly more efficient (up to eight orders of magnitude) for the lowest dust temperature in both the PDR and the core, while the formation of simple oxygen-bearing molecules (such as O_2) is much more efficient for the highest T_{dust} values in the core. Large temperature effects have also been found in the visual extinction threshold at which ice formation occurs, with variations of up to $\sim 5 \text{ mag}$ in the formation of water and CO_2 ices.

In this paper, we have also combined the steady-state Meijerink PDR code with the Nahoon time-dependent cloud code to realistically study the chemical evolution of several molecule families in the inner regions of different PDRs over 10^7 yr . The model results show that time dependence mainly affects the chemical evolution at intermediate and large visual extinctions, where long time-scales promote the destruction of oxygen-bearing molecules in low G_0 PDRs, and favour their formation and that of carbon-bearing molecules in high G_0 PDRs. Regarding solid species, we have found that CO_2 is a strongly time-dependent molecule, especially in low G_0 PDRs, as well as a late-forming ice compared to water, since the formation of its first ice monolayer occurs at $t \geq 10^5 \text{ yr}$ for $A_V \leq 10 \text{ mag}$. Formation of the first water ice monolayer occurs, however, at $t \sim 10^4 \text{ yr}$ for the same visual extinction range. This confirms the layered ice structure on dust grains (with H_2O in lower layers than CO_2) previously deduced by Cuppen et al. (2009) using CO, H_2CO , and CH_3OH ices.

Time evolution results also show that, at the edge of the cloud ($A_V \leq 1 \text{ mag}$), chemical steady state is reached at short times ($t \lesssim 10^5 \text{ yr}$) in any PDR type, being this time even shorter ($t < 10^4 \text{ yr}$) for high G_0 PDRs. By contrast, for larger visual extinctions, steady state strongly depends on the PDR characteristics. In high G_0 PDRs with densities $n \geq 10^5 \text{ cm}^{-3}$, chemistry reaches equilibrium at $t < 10^6 \text{ yr}$, while in low G_0 PDRs the time at which steady state is reached is $t > 10^6 \text{ yr}$.

The consideration of different types of PDRs in this study has also allowed us to shed light on the observed abundance enhancement of some COMs (e.g. CH_3CN) in PDR regions with respect to the cloud core (Gratier et al. 2013). From our theoretical results, we conclude that this enrichment is mainly due to the effect of increased density (which favours the formation of COM precursors, e.g. HCN in the case of CH_3CN) rather than to a direct UV radiation effect or the warming up of grain surfaces as previously suggested (Guzmán et al. 2014, Le Gal et al. 2017).

The results presented here, together with those from Esplugues et al. (2016), show how sensitive the chemistry of PDRs is to

the variation of their physical and chemical properties (density and intensity of radiation field), to the dust properties (dust temperature, type of grain substrate, efficiency of desorption), and to the evolutionary stage at each visual extinction. Our results also demonstrate how strongly coupled all these parameters are and the need of detailed observations of the physical and chemical structure of PDRs in order to put constraints on the chemical processes. This, together with the further exploration of the impact of other mechanisms associated to the PDR dynamics (e.g. turbulent diffusion or advection), as well as the implementation of more complex dust temperature treatments (e.g. mantle growth effects), will allow us to better reproduce observations and to gain a deeper understanding of the chemical evolution of PDRs.

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APPENDIX A

This Appendix lists (Tables A1–A6) all the surface reactions considered in the code. The new reactions included in this new version of the Meijerink PDR code are in italic. Binding energies are also shown (Table A3) for each species depending on the type of the grain substrate (bare or icy surface).

Table A1. Adsorption reactions.

Reaction ^a	Reaction
H → JH	CO ₂ → JCO ₂
H → JH _c	HCO → JHCO
H ₂ → JH ₂	H ₂ CO → JH ₂ CO
O → JO	CH ₃ O → JCH ₃ O
O ₂ → JO ₂	CH ₃ OH → JCH ₃ OH
O ₃ → JO ₃	N → JN
OH → JOH	N ₂ → JN ₂
H ₂ O → JH ₂ O	<i>S → JS</i>
HO ₂ → JHO ₂	<i>CH → JCH</i>
H ₂ O ₂ → JH ₂ O ₂	<i>CH₂ → JCH₂</i>
<i>C → JC</i>	<i>CH₃ → JCH₃</i>
CO → JCO	<i>CH₄ → JCH₄</i>

Note: ^a The expression Ji means solid *i*. The new reactions included in this version of the PDR code are in italic.

Table A2. Desorption reactions.

Reaction ^a	Reaction
JH → H	JCO ₂ → CO ₂
JH _c → H _c	JHCO → HCO
JH ₂ → H ₂	JH ₂ CO → H ₂ CO
JO → O	JCH ₃ O → CH ₃ O
JO ₂ → O ₂	JCH ₃ OH → CH ₃ OH
JO ₃ → O ₃	JN → N
JOH → OH	JN ₂ → N ₂
JH ₂ O → H ₂ O	<i>JS → S</i>
JHO ₂ → OH + O	<i>JCH → CH</i>
JH ₂ O ₂ → H ₂ O ₂	<i>JCH₂ → CH₂</i>
<i>JC → C</i>	<i>JCH₃ → CH₃</i>
JCO → CO	<i>JCH₄ → CH₄</i>

Note: ^a The expression Ji means solid *i*. The new reactions included in this version of the PDR code are in italic.

Table A3. Binding energies for the bare grain (E_b) and water ice (E_i) substrates.

Species	E_b (K)	E_i (K)	References
H	500	650	(1), (2)
H _c	10000	10000	(3)
H ₂	300	500	(4), (5)
O	1500	1420	(1), (6)
O ₂	1250	1160	(7)
O ₃	2200	2200	(8)
OH	4600	4600	(9)
H ₂ O	4800	5700	(10), (11)
HO ₂	4000	4000	(9)
H ₂ O ₂	6000	6000	(9)
CO	1200	1300	(12), (7)
CO ₂	3000	2670	(7), (13)
HCO	1600	1600	(14)
H ₂ CO	3700	3250	(15)
CH ₃ O	3700	3700	(16)
CH ₃ OH	3700	3700	(16)
N	720	720	(17)
N ₂	790	1140	(18)
S	1100	1100	(19)
C	800	800	(20)
CH	870	870	(21)
CH ₂	945	945	(21)
CH ₃	1017	1017	(21)
CH ₄	1090	1090	(22)

(1) Bergeron et al. (2008); (2) Al-Halabi & van Dishoeck (2007); (3) Cazaux & Tielens (2004); (4) Pirronello et al. (1997); (5) Amiaud et al. (2006); (6) Minissale (2014); (7) Noble et al. (2012a); (8) Borget et al. (2001); (9) Dulieu et al. (2013); (10) Sandford & Allamandola (1988); (11) Speedy et al. (1996); (12) Collings et al. (2003); (13) Karssemeijer & Cuppen (2014); (14) Garrod & Herbst (2006); (15) Noble et al. (2012b); (16) Collings et al. (2004); (17) Minissale et al. (2016); (18) Fuchs et al. (2006); (19) Aikawa (1997); (20) Tielens & Allamandola (1987); (21) Taquet, Charnley & Sipilä (2014); (22) Herrero et al. (2010).

Table A4. Reactions on grain surfaces.

Reaction	δ_{bare}^a	δ_{ice}^a	ϵ^b
JH + JH \rightarrow H ₂	9.630E-1	9.640E-2	0
JH + JH \rightarrow JH ₂	3.700E-2	9.036E-1	0
JH + JO \rightarrow OH	3.875E-1	3.880E-2	0
JH + JO \rightarrow JOH	6.125E-1	9.612E-1	0
JH + JOH \rightarrow H ₂ O	2.677E-1	2.680E-2	0
JH + JOH \rightarrow JH ₂ O	7.323E-1	9.732E-1	0
JH + JHO ₂ \rightarrow H ₂ O ₂	4.600E-3	5.000E-4	0
JH + JHO ₂ \rightarrow JH ₂ O ₂	9.954E-1	9.995E-1	0
JH + JO ₂ \rightarrow HO ₂	1.380E-2	1.400E-3	0
JH + JO ₂ \rightarrow JHO ₂	9.862E-1	9.986E-1	0
JH + JCO \rightarrow HCO	6.700E-3	7.000E-4	2000
JH + JCO \rightarrow JHCO	9.933E-1	9.993E-1	2000
JH + JHCO \rightarrow H ₂ CO	6.610E-2	6.700E-3	0
JH + JHCO \rightarrow JH ₂ CO	9.339E-1	9.933E-1	0
JH + JH ₂ CO \rightarrow CH ₃ O	1.000E-4	1.000E-4	2000
JH + JH ₂ CO \rightarrow JCH ₃ O	9.999E-1	9.999E-1	2000
JH + JCH ₃ O \rightarrow CH ₃ OH	2.350E-2	2.400E-3	0
JH + JCH ₃ O \rightarrow JCH ₃ OH	9.765E-1	9.976E-1	0
JH + JC \rightarrow CH	8.212E-1	8.220E-2	0
JH + JC \rightarrow JCH	1.788E-1	9.178E-1	0
JH + JCH \rightarrow CH ₂	7.668E-1	7.670E-2	0
JH + JCH \rightarrow JCH ₂	2.332E-1	9.233E-1	0
JH + JCH ₂ \rightarrow CH ₃	6.937E-1	6.940E-2	0
JH + JCH ₂ \rightarrow JCH ₃	3.063E-1	9.306E-1	0
JH + JCH ₃ \rightarrow CH ₄	5.886E-1	5.890E-2	0
JH + JCH ₃ \rightarrow JCH ₄	4.114E-1	9.411E-1	0
JO + JO \rightarrow O ₂	6.884E-1	6.890E-2	0
JO + JO \rightarrow JO ₂	3.116E-1	9.311E-1	0
JO + JC \rightarrow CO	8.659E-1	8.660E-2	0
JO + JC \rightarrow JCO	1.341E-1	9.134E-1	0
JO + JCO \rightarrow CO ₂	1.403E-1	1.400E-2	650
JO + JCO \rightarrow JCO ₂	8.597E-1	9.860E-1	650
JO + JO ₂ \rightarrow O ₃	3.000E-4	1.000E-4	0
JO + JO ₂ \rightarrow JO ₃	9.997E-1	9.999E-1	0
JO + JCH ₃ \rightarrow JCH ₃ O	1.000E + 0	1.000E + 0	0
JOH + JOH \rightarrow H ₂ O ₂	2.000E-4	1.000E-4	0
JOH + JOH \rightarrow JH ₂ O ₂	9.998E-1	9.999E-1	0
JOH + JCH ₂ \rightarrow JCH ₃ O	1.000E + 0	1.000E + 0	0
JN + JN \rightarrow N ₂	8.977E-1	8.980E-2	0
JN + JN \rightarrow JN ₂	1.023E-1	9.102E-1	0
JH + JO ₃ \rightarrow OH + O ₂	8.020E-2	8.100E-3	480
JH + JO ₃ \rightarrow OH + JO ₂	0.000E + 0	0.000E + 0	480
JH + JO ₃ \rightarrow JOH + O ₂	2.346E-1	2.340E-2	480
JH + JO ₃ \rightarrow JOH + JO ₂	6.852E-1	9.685E-1	480
JH + JH ₂ O \rightarrow OH + H ₂	0.000E + 0	0.000E + 0	9600
JH + JH ₂ O \rightarrow OH + JH ₂	0.000E + 0	0.000E + 0	9600
JH + JH ₂ O \rightarrow JOH + H ₂	0.000E + 0	0.000E + 0	9600
JH + JH ₂ O \rightarrow JOH + JH ₂	1.000E + 0	1.000E + 0	9600
JH + JHO ₂ \rightarrow OH + OH	3.400E-3	4.000E-4	0
JH + JHO ₂ \rightarrow OH + JOH	0.000E + 0	0.000E + 0	0
JH + JHO ₂ \rightarrow JOH + OH	0.000E + 0	0.000E + 0	0
JH + JHO ₂ \rightarrow JOH + JOH	9.966E-1	9.996E-1	0
JH + JH ₂ O ₂ \rightarrow H ₂ O + OH	2.120E-2	2.100E-3	1000
JH + JH ₂ O ₂ \rightarrow JH ₂ O + OH	7.200E-3	8.000E-4	1000
JH + JH ₂ O ₂ \rightarrow H ₂ O + JOH	0.000E + 0	0.000E + 0	1000
JH + JH ₂ O ₂ \rightarrow JH ₂ O + JOH	9.716E-1	9.971E-1	1000
JH + JHCO \rightarrow CO + H ₂	4.347E-1	4.360E-2	0
JH + JHCO \rightarrow CO + JH ₂	0.000E + 0	0.000E + 0	0
JH + JHCO \rightarrow JCO + H ₂	4.827E-1	4.820E-2	0
JH + JHCO \rightarrow JCO + JH ₂	8.260E-2	9.082E-1	0
JH + JH ₂ CO \rightarrow HCO + H ₂	2.000E-4	1.000E-4	2200
JH + JH ₂ CO \rightarrow HCO + JH ₂	0.000E + 0	0.000E + 0	2200
JH + JH ₂ CO \rightarrow JHCO + H ₂	5.050E-1	5.050E-2	2200
JH + JH ₂ CO \rightarrow JHCO + JH ₂	4.948E-1	9.494E-1	2200

Table A4 – continued

Reaction	δ_{bare}^a	δ_{ice}^a	ϵ^b
JH + JCH ₃ O \rightarrow H ₂ CO + H ₂	1.260E-2	1.400E-3	150
JH + JCH ₃ O \rightarrow H ₂ CO + JH ₂	0.000E + 0	0.000E + 0	150
JH + JCH ₃ O \rightarrow JH ₂ CO + H ₂	8.596E-1	8.590E-2	150
JH + JCH ₃ O \rightarrow JH ₂ CO + JH ₂	1.278E-1	9.127E-1	150
JH + JH ₄ CO \rightarrow CH ₃ O + H ₂	0.000E + 0	0.000E + 0	3200
JH + JH ₄ CO \rightarrow CH ₃ O + JH ₂	0.000E + 0	0.000E + 0	3200
JH + JH ₄ CO \rightarrow JCH ₃ O + H ₂	0.000E + 0	0.000E + 0	3200
JH + JH ₄ CO \rightarrow JCH ₃ O + JH ₂	1.000E + 0	1.000E + 0	3200
JH + JCO ₂ \rightarrow CO + OH	0.000E + 0	0.000E + 0	10000
JH + JCO ₂ \rightarrow CO + JOH	0.000E + 0	0.000E + 0	10000
JH + JCO ₂ \rightarrow JCO + OH	0.000E + 0	0.000E + 0	10000
JH + JCO ₂ \rightarrow JCO + JOH	1.000E + 0	1.000E + 0	10000
JH + JCH \rightarrow C + H ₂	3.915E-1	3.915E-2	0
JH + JCH \rightarrow C + JH ₂	0.000E + 0	0.000E + 0	0
JH + JCH \rightarrow JC + H ₂	3.859E-1	3.865E-2	0
JH + JCH \rightarrow JC + JH ₂	2.226E-1	9.222E-1	0
JH + JCH ₂ \rightarrow CH + H ₂	8.000E-5	4.800E-5	0
JH + JCH ₂ \rightarrow CH + JH ₂	0.000E + 0	0.000E + 0	0
JH + JCH ₂ \rightarrow JCH + H ₂	4.452E-2	4.452E-3	0
JH + JCH ₂ \rightarrow JCH + JH ₂	9.554E-1	9.955E-1	0
JH + JCH ₃ \rightarrow CH ₂ + H ₂	0.000E + 0	0.000E + 0	0
JH + JCH ₃ \rightarrow CH ₂ + JH ₂	0.000E + 0	0.000E + 0	0
JH + JCH ₃ \rightarrow JCH ₂ + H ₂	0.000E + 0	0.000E + 0	0
JH + JCH ₃ \rightarrow JCH ₂ + JH ₂	1.000E + 0	1.000E + 0	0
JH + JCH ₄ \rightarrow CH ₃ + H ₂	0.000E + 0	0.000E + 0	0
JH + JCH ₄ \rightarrow CH ₃ + JH ₂	0.000E + 0	0.000E + 0	0
JH + JCH ₄ \rightarrow JCH ₃ + H ₂	0.000E + 0	0.000E + 0	0
JH + JCH ₄ \rightarrow JCH ₃ + JH ₂	1.000E + 0	1.000E + 0	0
JO + JO ₃ \rightarrow O ₂ + O ₂	3.872E-1	3.880E-2	2500
JO + JO ₃ \rightarrow O ₂ + JO ₂	0.000E + 0	0.000E + 0	2500
JO + JO ₃ \rightarrow JO ₂ + O ₂	0.000E + 0	0.000E + 0	2500
JO + JO ₃ \rightarrow JO ₂ + JO ₂	6.128E-1	9.612E-1	2500
JO + JOH \rightarrow O ₂ + H	1.890E-2	2.000E-3	0
JO + JOH \rightarrow O ₂ + JH	0.000E + 0	0.000E + 0	0
JO + JOH \rightarrow JO ₂ + H	5.264E-1	5.260E-2	0
JO + JOH \rightarrow JO ₂ + JH	4.547E-1	9.454E-1	0
JO + JHO ₂ \rightarrow O ₂ + OH	2.190E-2	2.300E-3	0
JO + JHO ₂ \rightarrow O ₂ + JOH	1.516E-1	1.510E-2	0
JO + JHO ₂ \rightarrow JO ₂ + OH	0.000E + 0	0.000E + 0	0
JO + JHO ₂ \rightarrow JO ₂ + JOH	8.265E-1	9.826E-1	0
JO + JHCO \rightarrow CO ₂ + H	5.110E-2	5.200E-3	0
JO + JHCO \rightarrow CO ₂ + JH	0.000E + 0	0.000E + 0	0
JO + JHCO \rightarrow JCO ₂ + H	8.348E-1	8.340E-2	0
JO + JHCO \rightarrow JCO ₂ + JH	1.141E-1	9.114E-1	0
JO + JH ₂ CO \rightarrow CO ₂ + H ₂	3.680E-2	3.700E-3	335
JO + JH ₂ CO \rightarrow CO ₂ + JH ₂	0.000E + 0	0.000E + 0	335
JO + JH ₂ CO \rightarrow JCO ₂ + H ₂	8.901E-1	8.900E-2	335
JO + JH ₂ CO \rightarrow JCO ₂ + JH ₂	7.310E-2	9.073E-1	335
JOH + JH ₂ \rightarrow H ₂ O + H	0.000E + 0	0.000E + 0	2100
JOH + JH ₂ \rightarrow H ₂ O + JH	0.000E + 0	0.000E + 0	2100
JOH + JH ₂ \rightarrow JH ₂ O + H	4.052E-1	4.060E-2	2100
JOH + JH ₂ \rightarrow JH ₂ O + JH	5.948E-1	9.594E-1	2100
JOH + JCO \rightarrow CO ₂ + H	1.000E-4	6.000E-5	400
JOH + JCO \rightarrow CO ₂ + JH	0.000E + 0	0.000E + 0	400
JOH + JCO \rightarrow JCO ₂ + H	5.794E-1	5.794E-2	400
JOH + JCO \rightarrow JCO ₂ + JH	4.205E-1	9.420E-1	400
JOH + JHCO \rightarrow CO ₂ + H ₂	2.577E-2	2.640E-3	0
JOH + JHCO \rightarrow CO	0.000E-0	0.000E-0	0
JOH + JHCO \rightarrow JCO ₂ + H ₂	8.936E-1	8.936E-2	0
JOH + JHCO \rightarrow JCO ₂ + JH ₂	8.063E-2	9.080E-1	0
JOH + JCH ₃ OH \rightarrow JCH ₃ O + JH ₂ O	1.000E + 0	1.000E + 0	5000
JHO ₂ + JH ₂ \rightarrow H ₂ O ₂ + H	0.000E + 0	0.000E + 0	5000
JHO ₂ + JH ₂ \rightarrow H ₂ O ₂ + JH	0.000E + 0	0.000E + 0	5000

Table A4 – *continued*

Reaction	δ_{bare}^a	δ_{ice}^a	ϵ^b
$\text{JHO}_2 + \text{JH}_2 \rightarrow \text{JH}_2\text{O}_2 + \text{H}$	0.000E + 0	0.000E + 0	5000
$\text{JHO}_2 + \text{JH}_2 \rightarrow \text{JH}_2\text{O}_2 + \text{JH}$	1.000E + 0	1.000E + 0	5000

The expression Ji means solid i .

^aThe parameters δ_{bare} and δ_{ice} indicate the probabilities of desorption upon reaction for bare and icy substrates, respectively.

^bThe parameter ϵ indicates the activation barrier for each reaction. The new reactions included in this version of the PDR code are in *italic*.

Table A5. Photoreactions on dust grains.

Reaction ^a	α_i^b (s ⁻¹)	ξ_i^b
<i>JCO + Photon</i> \rightarrow <i>JC + JO</i>	2.59×10^{-10}	3.53
<i>JH}_2 + \text{Photon}</i> \rightarrow <i>JH + JH</i>	8.00×10^{-10}	2.20
<i>JO}_2 + \text{Photon}</i> \rightarrow <i>JO + JO</i>	7.90×10^{-10}	2.13
<i>JOH + Photon</i> \rightarrow <i>JH + JO</i>	3.90×10^{-10}	2.24
<i>JCO}_2 + \text{Photon}</i> \rightarrow <i>JO + JCO</i>	8.90×10^{-10}	3.00
<i>JH}_2\text{O} + \text{Photon}</i> \rightarrow <i>JH + JOH</i>	8.00×10^{-10}	2.20
<i>JHCO + Photon</i> \rightarrow <i>JH + JCO</i>	1.10×10^{-09}	1.09
<i>JH}_2\text{CO} + \text{Photon}</i> \rightarrow <i>JH + JHCO</i>	5.87×10^{-10}	0.53
<i>JCH}_3\text{O} + \text{Photon}</i> \rightarrow <i>JH + JH}_2\text{CO}</i>	5.87×10^{-10}	0.53
<i>JCH}_3\text{OH} + \text{Photon}</i> \rightarrow <i>JH + JCH}_3\text{O}</i>	5.87×10^{-10}	0.53
<i>JN}_2 + \text{Photon}</i> \rightarrow <i>JN + JN</i>	2.30×10^{-10}	3.88
<i>JHO}_2 + \text{Photon}</i> \rightarrow <i>JO + JOH</i>	3.28×10^{-10}	1.63
<i>JHO}_2 + \text{Photon}</i> \rightarrow <i>JO}_2 + JH</i>	3.28×10^{-10}	1.63
<i>JH}_2\text{O}_2 + \text{Photon}</i> \rightarrow <i>JOH + JOH</i>	8.30×10^{-10}	1.80
<i>JO}_3 + \text{Photon}</i> \rightarrow <i>JO}_2 + JO</i>	3.30×10^{-10}	1.40
<i>JCH + Photon</i> \rightarrow <i>C + H</i>	9.20×10^{-10}	1.72
<i>JCH}_2 + \text{Photon}</i> \rightarrow <i>H + CH</i>	5.80×10^{-10}	2.02
<i>JCH}_3 + \text{Photon}</i> \rightarrow <i>H + CH}_2</i>	1.35×10^{-10}	2.27
<i>JCH}_4 + \text{Photon}</i> \rightarrow <i>H}_2 + CH}_2</i>	7.20×10^{-10}	2.59
<i>JCO + Photon</i> \rightarrow <i>CO</i>	3.67×10^{-10}	2.54
<i>JH}_2\text{O} + \text{Photon}</i> \rightarrow <i>H}_2\text{O}</i>	3.67×10^{-11}	2.20
<i>JHCO + Photon</i> \rightarrow <i>HCO</i>	3.67×10^{-11}	0.53
<i>JH}_2\text{CO} + \text{Photon}</i> \rightarrow <i>H}_2\text{CO}</i>	3.67×10^{-11}	0.53
<i>JCH}_3\text{O} + \text{Photon}</i> \rightarrow <i>CH}_3\text{O}</i>	3.67×10^{-11}	0.53
<i>JCH}_3\text{OH} + \text{Photon}</i> \rightarrow <i>CH}_3\text{OH}</i>	5.00×10^{-13}	0.53
<i>JCH}_3\text{OH} + \text{Photon}</i> \rightarrow <i>CH}_3\text{O} + \text{H}</i>	5.00×10^{-13}	0.53

^aThe expression Ji means solid i .

^bValues for α_i and ξ_i (dimensionless) are taken from KIDA. The new reactions included in this version of the PDR code are in *italic*.

Table A6. Cosmic ray reactions and photo processes induced by cosmic rays.

Reaction ^a	κ_{CR}^b (s ⁻¹)
<i>JH}_2 + \text{CR}</i> \rightarrow <i>JH + JH</i>	5.00×10^{-17}
<i>JO}_2 + \text{CRP}</i> \rightarrow <i>JO + JO</i>	3.75×10^{-14}
<i>JOH + CRP</i> \rightarrow <i>JH + JO</i>	2.55×10^{-14}
<i>JCO}_2 + \text{CRP}</i> \rightarrow <i>JO + JCO</i>	8.55×10^{-14}
<i>JH}_2\text{O} + \text{CRP}</i> \rightarrow <i>JH + JOH</i>	4.85×10^{-14}
<i>JHCO + CRP</i> \rightarrow <i>JH + JCO</i>	2.11×10^{-14}
<i>JH}_2\text{CO} + \text{CRP}</i> \rightarrow <i>JH + JHCO</i>	2.11×10^{-14}
<i>JCH}_3\text{O} + \text{CRP}</i> \rightarrow <i>JH + JH}_2\text{CO}</i>	2.11×10^{-14}
<i>JCH}_3\text{OH} + \text{CRP}</i> \rightarrow <i>JH + JCH}_3\text{O}</i>	2.11×10^{-14}
<i>JN}_2 + \text{CRP}</i> \rightarrow <i>JN + JN</i>	2.50×10^{-16}
<i>JHO}_2 + \text{CRP}</i> \rightarrow <i>JO + JOH</i>	3.75×10^{-14}
<i>JHO}_2 + \text{CRP}</i> \rightarrow <i>JH + JO}_2</i>	3.75×10^{-14}
<i>JH}_2\text{O}_2 + \text{CRP}</i> \rightarrow <i>JOH + JOH</i>	7.50×10^{-14}
<i>JO}_3 + \text{CRP}</i> \rightarrow <i>JO}_2 + JO</i>	3.75×10^{-14}
<i>JCO + CRP</i> \rightarrow <i>CO</i>	1.08×10^{-14}
<i>JH}_2\text{O} + \text{CRP}</i> \rightarrow <i>H}_2\text{O}</i>	1.08×10^{-14}
<i>JH}_2\text{CO} + \text{CRP}</i> \rightarrow <i>H}_2\text{CO}</i>	1.08×10^{-14}
<i>JCH}_3\text{OH} + \text{CRP}</i> \rightarrow <i>CH}_3\text{OH}</i>	1.08×10^{-14}
<i>JCH + CRP</i> \rightarrow <i>C + H</i>	3.65×10^{-14}
<i>JCH}_3 + \text{CRP}</i> \rightarrow <i>H + CH}_2</i>	2.50×10^{-14}
<i>JCH}_4 + \text{CRP}</i> \rightarrow <i>H}_2 + CH}_2</i>	1.17×10^{-13}

^aThe expression Ji means solid i .

^bValues for the cosmic ray rate coefficient, κ_{CR} , are taken from KIDA. The new reactions included in this version of the PDR code are in *italic*.

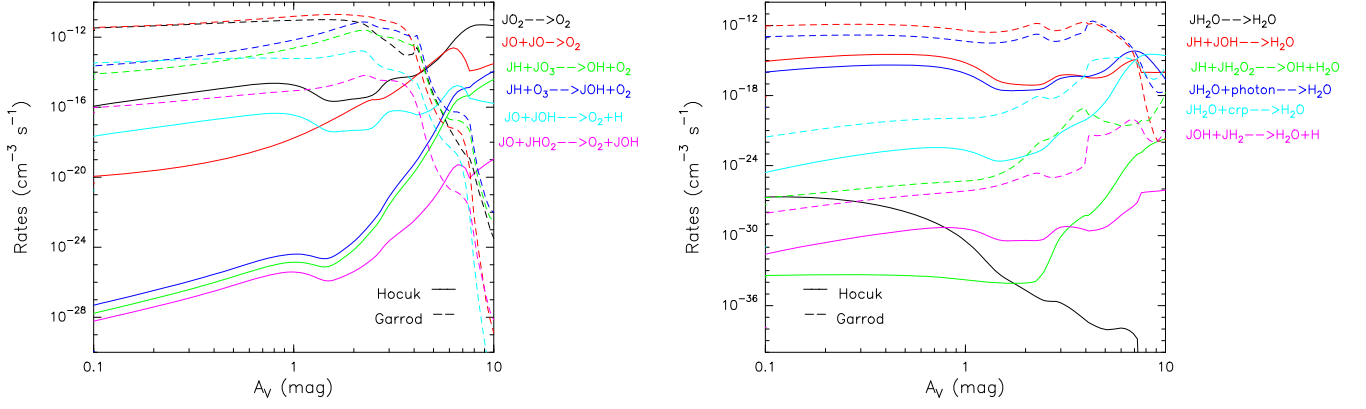


Figure A1. Rates for surface reactions forming O_2 gas (left) and H_2O gas (right) for Model 2 ($G_0 = 10^4$ and $n = 10^5 cm^{-3}$) using T_{dust} from Hocuk et al. (2017) (solid lines) and from Garrod & Pauly (2011) (dashed lines). JX means solid X.

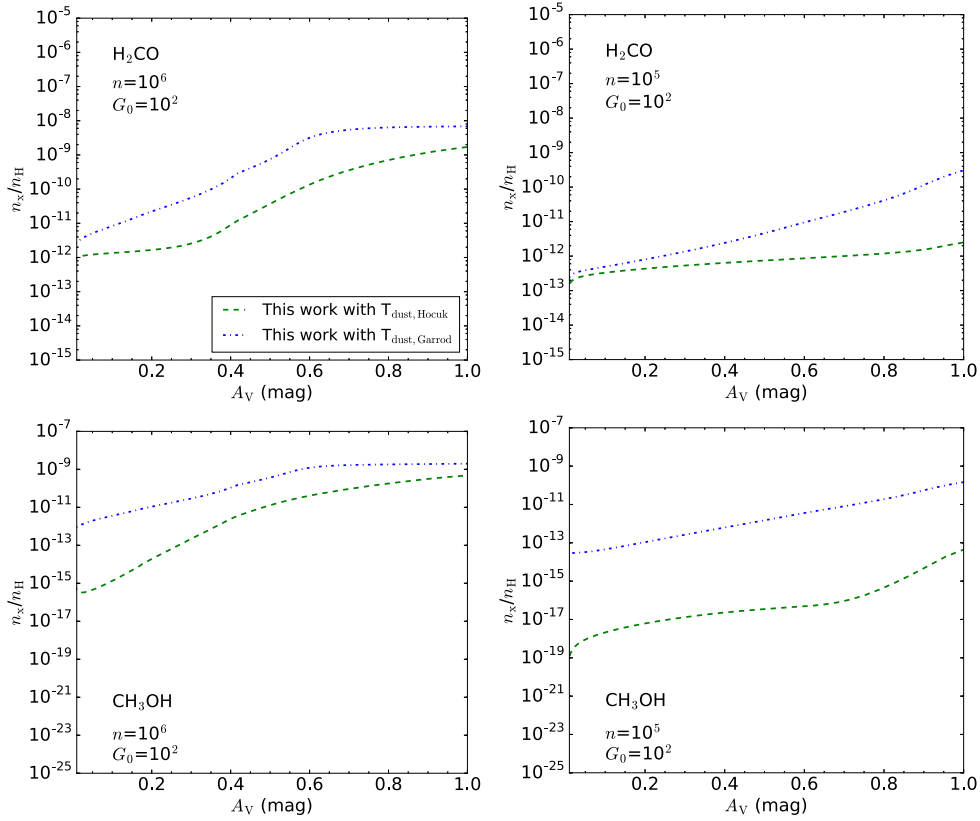


Figure A2. Abundances of H_2CO and CH_3OH obtained with the updated Meijerink PDR code using T_{dust} from Hocuk et al. (2017) (green dashed lines) and from Garrod & Pauly (2011) (blue dotted lines) for a model with $n = 10^6 cm^{-3}$ and $G_0 = 10^2$ (left column), and with $n = 10^5 cm^{-3}$ and $G_0 = 10^2$ (right column).

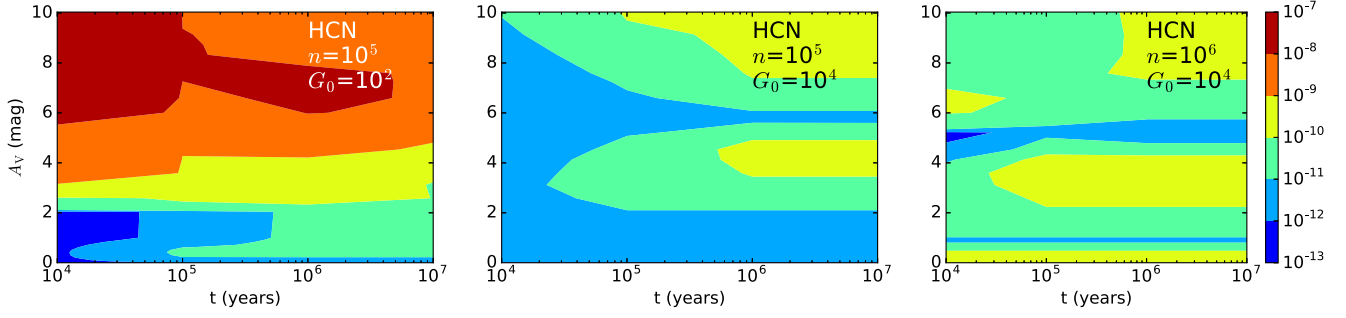


Figure A3. Contour maps with the abundances of HCN for Models 1 (left-hand panel), 2 (middle panel), and 3 (right-hand panel) as a function of time (x-axis) and visual extinction (y-axis).

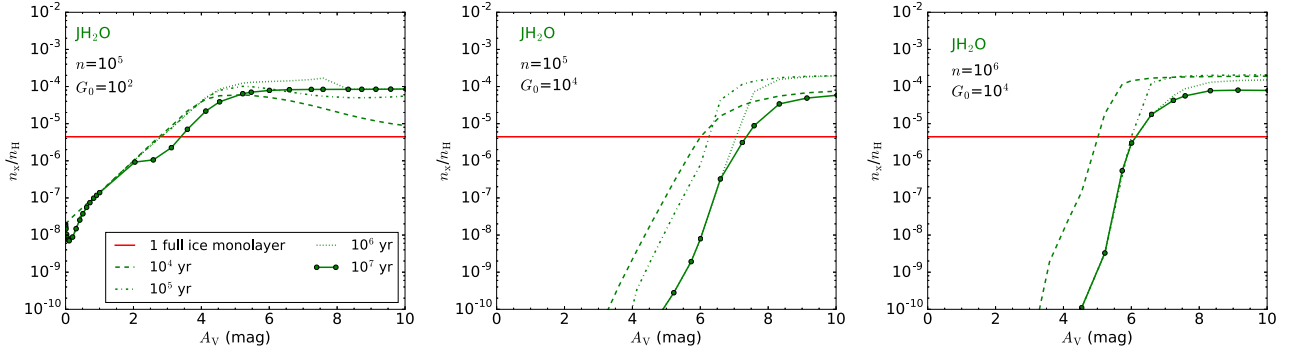


Figure A4. Fractional abundances of solid H_2O for Models 1 (left-hand panel), 2 (middle panel), and 3 (right-hand panel) as a function of the visual extinction at different time-scales: 10^4 yr (dashed line), 10^5 yr (dash-dotted line), 10^6 yr (dotted line), and 10^7 yr (solid-dotted line). J_i means solid i . The red line represents the number of possible adsorption sites on grain surfaces per cm^2 .

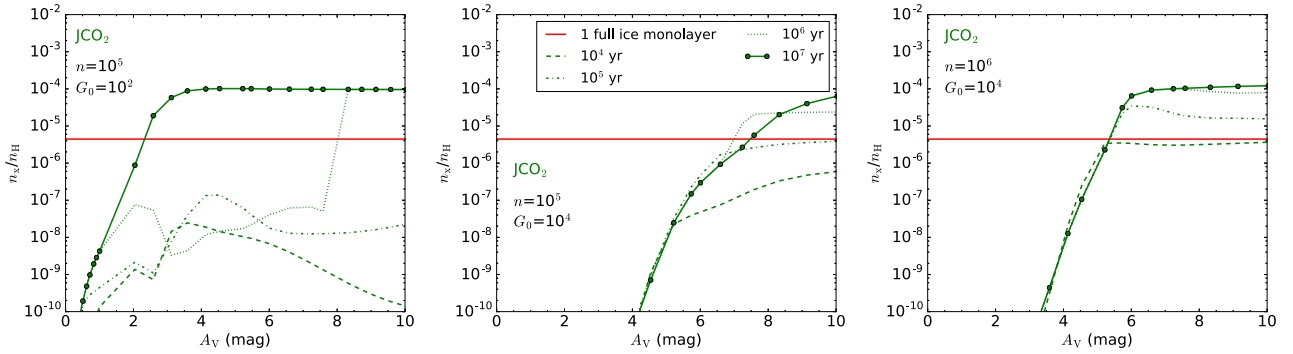


Figure A5. Fractional abundances of solid CO_2 for Models 1 (left-hand panel), 2 (middle panel), and 3 (right-hand panel) as a function of the visual extinction at different time-scales: 10^4 yr (dashed line), 10^5 yr (dash-dotted line), 10^6 yr (dotted line), and 10^7 yr (solid-dotted line). J_i means solid i . The red line represents the number of possible adsorption sites on grain surfaces per cm^2 .

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